

PATENT COOPERATION TREATY

PCT

NOTIFICATION OF ELECTION

(PCT Rule 61.2)

From the INTERNATIONAL BUREAU

To:

Assistant Commissioner for Patents
United States Patent and Trademark
Office
Box PCT
Washington, D.C. 20231
ÉTATS-UNIS D'AMÉRIQUE

in its capacity as elected Office

Date of mailing (day/month/year) 06 December 1999 (06.12.99)	Applicant's or agent's file reference FCCC 98-02
International application No. PCT/US99/06644	Priority date (day/month/year) 27 March 1998 (27.03.98)
International filing date (day/month/year) 26 March 1999 (26.03.99)	
Applicant KRUH, Gary et al	

1. The designated Office is hereby notified of its election made:

☒ in the demand filed with the International Preliminary Examining Authority on:

22 October 1999 (22.10.99)

☐ in a notice effecting later election filed with the International Bureau on:2. The election ☒ was☐ was not

made before the expiration of 19 months from the priority date or, where Rule 32 applies, within the time limit under Rule 32.2(b).

<p>The International Bureau of WIPO 34, chemin des Colombettes 1211 Geneva 20, Switzerland</p> <p>Facsimile No.: (41-22) 740.14.35</p>	<p>Authorized officer Olivia RANAIVOJAONA</p> <p>Telephone No.: (41-22) 338.83.38</p>
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PATENT COOPERATION TREATY

WO 99/49735
PCT/US99/06644

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NOTICE INFORMING THE APPLICANT OF THE COMMUNICATION OF THE INTERNATIONAL APPLICATION TO THE DESIGNATED OFFICES

(PCT Rule 47.1(c), first sentence)

From the INTERNATIONAL BUREAU

To:

RIGAUT, Kathleen, D.
Dann, Dorfman, Herrell and Skillman
Suite 720
1601 Market Street
Philadelphia, PA 19103
ÉTATS-UNIS D'AMÉRIQUE

Date of mailing (day/month/year) 07 October 1999 (07.10.99)		IMPORTANT NOTICE	
Applicant's or agent's file reference FCCC 98-02			
International application No. PCT/US99/06644	International filing date (day/month/year) 26 March 1999 (26.03.99)	Priority date (day/month/year) 27 March 1998 (27.03.98)	
Applicant FOX CHASE CANCER CENTER et al			

1. Notice is hereby given that the International Bureau has communicated, as provided in Article 20, the international application to the following designated Offices on the date indicated above as the date of mailing of this Notice:
AU,EP,JP,US

In accordance with Rule 47.1(c), third sentence, those Offices will accept the present Notice as conclusive evidence that the communication of the international application has duly taken place on the date of mailing indicated above and no copy of the international application is required to be furnished by the applicant to the designated Office(s).

2. The following designated Offices have waived the requirement for such a communication at this time:
CA

The communication will be made to those Offices only upon their request. Furthermore, those Offices do not require the applicant to furnish a copy of the international application (Rule 49.1(a-bis)).

3. Enclosed with this Notice is a copy of the international application as published by the International Bureau on
07 October 1999 (07.10.99) under No. WO 99/49735

REMINDER REGARDING CHAPTER II (Article 31(2)(a) and Rule 54.2)

If the applicant wishes to postpone entry into the national phase until 30 months (or later in some Offices) from the priority date, a **demand for international preliminary examination** must be filed with the competent International Preliminary Examining Authority before the expiration of 19 months from the priority date.

It is the applicant's sole responsibility to monitor the 19-month time limit.

Note that only an applicant who is a national or resident of a PCT Contracting State which is bound by Chapter II has the right to file a demand for international preliminary examination.

REMINDER REGARDING ENTRY INTO THE NATIONAL PHASE (Article 22 or 39(1))

If the applicant wishes to proceed with the international application in the **national phase**, he must, within 20 months or 30 months, or later in some Offices, perform the acts referred to therein before each designated or elected Office.

For further important information on the time limits and acts to be performed for entering the national phase, see the Annex to Form PCT/IB/301 (Notification of Receipt of Record Copy) and Volume II of the PCT Applicant's Guide.

<p>The International Bureau of WIPO 34, chemin des Colombettes 1211 Geneva 20, Switzerland</p> <p>Facsimile No. (41-22) 740.14.35</p>	<p>Authorized officer J. Zahra</p> <p>Telephone No. (41-22) 338.83.38</p>
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REC'D 03 JUL 2000

WIPO

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INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Article 36 and Rule 70)

Applicant's or agent's file reference FCCC 98-02	FOR FURTHER ACTION See Notification of Transmittal of International Preliminary Examination Report (Form PCT/IPEA/416)	
International application No. PCT/US99/06644	International filing date (day/month/year) 26 MARCH 1999	Priority date (day/month/year) 27 MARCH 1998
International Patent Classification (IPC) or national classification and IPC Please See Supplemental Sheet.		
Applicant FOX CHASE CANCER CENTER		

1. This international preliminary examination report has been prepared by this International Preliminary Examining Authority and is transmitted to the applicant according to Article 36.

2. This REPORT consists of a total of 5 sheets.

☐ This report is also accompanied by ANNEXES, i.e., sheets of the description, claims and/or drawings which have been amended and are the basis for this report and/or sheets containing rectifications made before this Authority. (see Rule 70.16 and Section 607 of the Administrative Instructions under the PCT).

These annexes consist of a total of 0 sheets.

3. This report contains indications relating to the following items:

- I ☒ Basis of the report
- II ☐ Priority
- III ☐ Non-establishment of report with regard to novelty, inventive step or industrial applicability
- IV ☐ Lack of unity of invention
- V ☒ Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement
- VI ☐ Certain documents cited
- VII ☐ Certain defects in the international application
- VIII ☒ Certain observations on the international application

Date of submission of the demand 22 OCTOBER 1999	Date of completion of this report 12 JUNE 2000
Name and mailing address of the IPEA/US Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231	Authorized officer <i>Shin-Lin Chen</i> SHIN-LIN CHEN
Facsimile No. (703) 305-3230	Telephone No. (703) 308-0196

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.

PCT/US99/06644

I. Basis of the report

1. With regard to the elements of the international application:*

- ☒ the international application as originally filed
- ☒ the description:
pages 1-62 , as originally filed
pages NONE , filed with the demand
pages NONE , filed with the letter of _____
- ☒ the claims:
pages 63-71 , as originally filed
pages NONE , as amended (together with any statement) under Article 19
pages NONE , filed with the demand
pages NONE , filed with the letter of _____
- ☒ the drawings:
pages 1-56 , as originally filed
pages NONE , filed with the demand
pages NONE , filed with the letter of _____
- ☒ the sequence listing part of the description:
pages 1-19 , as originally filed
pages NONE , filed with the demand
pages NONE , filed with the letter of _____

2. With regard to the language, all the elements marked above were available or furnished to this Authority in the language in which the international application was filed, unless otherwise indicated under this item.

These elements were available or furnished to this Authority in the following language _____ which is:

- ☐ the language of a translation furnished for the purposes of international search (under Rule 23.1(b)).
- ☐ the language of publication of the international application (under Rule 48.3(b)).
- ☐ the language of the translation furnished for the purposes of international preliminary examination (under Rules 55.2 and/or 55.3).

3. With regard to any nucleotide and/or amino acid sequence disclosed in the international application, the international preliminary examination was carried out on the basis of the sequence listing:

- ☒ contained in the international application in printed form.
- ☒ filed together with the international application in computer readable form.
- ☐ furnished subsequently to this Authority in written form.
- ☐ furnished subsequently to this Authority in computer readable form.
- ☐ The statement that the subsequently furnished written sequence listing does not go beyond the disclosure in the international application as filed has been furnished.
- ☐ The statement that the information recorded in computer readable form is identical to the written sequence listing has been furnished.

4. ☒ The amendments have resulted in the cancellation of:

- ☒ the description, pages NONE
- ☒ the claims, Nos. NONE
- ☒ the drawings, sheets/fig NONE

5. ☒ This report has been drawn as if (some of) the amendments had not been made, since they have been considered to go beyond the disclosure as filed, as indicated in the Supplemental Box (Rule 70.2(c)).**

* Replacement sheets which have been furnished to the receiving Office in response to an invitation under Article 14 are referred to in this report as "originally filed" and are not annexed to this report since they do not contain amendments (Rules 70.16 and 70.17).

**Any replacement sheet containing such amendments must be referred to under item 1 and annexed to this report.

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.

PCT/US99/06644

V. Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement**1. statement**

Novelty (N)	Claims <u>1-20, 23-32, 34-43, 45-59</u>	YES
	Claims <u>21, 22, 33, 44</u>	NO
Inventive Step (IS)	Claims <u>1-20, 23-32, 34-43, 45-59</u>	YES
	Claims <u>21, 22, 33, 44</u>	NO
Industrial Applicability (IA)	Claims <u>1-59</u>	YES
	Claims <u>NONE</u>	NO

2. citations and explanations (Rule 70.7)

Claims 21, 22, 33 and 44 lack novelty under PCT Article 33(2) as being anticipated by GenBank Accession Nos. U66687, D77412, U66674 and R97754, respectively.

Because U66687 has nucleotide sequence 97.4% identical to base 4064 to 4808 of SEQ ID NO. 3 encoding amino acids of SEQ ID NO. 4, D77412 has nucleotide sequence 82.2% identical to base 134 to 408 of SEQ ID NO. 1 encoding the amino acids of SEQ ID NO. 2, U66674 contains nucleotide sequence 97.7% identical to base 1946 to 3134 of SEQ ID No. 5 encoding the amino acids of SEQ ID NO. 6, and R97754 contains nucleotide sequence 98.2% identical to base 4 to 221 of SEQ ID NO. 7 encoding the amino acids of SEQ ID NO. 8. Thus claims 21, 22, 33 and 44 are clearly anticipated by GenBank Accession Nos. U66687, D77412, U66674 and R97754, respectively.

Since claims 21, 22, 33 and 44 all lack novelty, so they also lack inventive step. On the other hand, claims 1-20, 23-32, 34-43 and 45-59 all have inventive step and novelty.

----- NEW CITATIONS -----

VIII. Certain observations on the international application

The following observations on the clarity of the claims, description, and drawings or on the question whether the claims are fully supported by the description, are made:

Claim 44 is objected to under PCT Rule 66.2(a)(v) as lacking clarity under PCT Article 6 because the claim is indefinite for the following reason(s): SEQ ID NO. 7 represents nucleotide sequence instead of amino acid sequence.

Claims 53-55 are objected to under PCT Rule 66.2(a)(v) as lacking clarity under PCT Article 6 because the claim are indefinite for the following reason(s): There is no antecedent basis for the term "said mouse" in claims 53-55.

The description is objected to under PCT Rule 66.2(a)(v) as lacking clarity under PCT Article 5 because it fails to adequately enable practice of the claimed invention because: Claims 48-52 are drawn to a host cell comprising the nucleotide sequence of SEQ ID No. 1, 3, 5 or 7 and a host animal comprising said nucleotide sequence *in vivo*. Claims 53-55 are drawn to a host animal, such as a transgenic mouse, harboring a homozygous null mutation in its endogenous MOAT gene. It was unpredictable at the time of the invention in making a transgenic animal harboring a transgene under the control of a promoter. One skilled in the art would not be able to predict the phenotype of the transgenic animal produced. The vector used, the coding sequence, the non-coding sequence, the promoter and the integration site of the transgene in the genome of the host cells are all important factors in contributing to the resulting phenotypes of a transgenic animal. The specification of the present application fails to provide adequate guidance for making a transgenic animal via embryonic stem cells except using mouse embryonic stem cells. Thus, it would have required a skilled artisan to engage in undue experimentation to practice the claimed invention. The description of the present application neither enables making a transgenic animal harboring any transgene, nor enables making a host cell derived from said transgenic animal.

Claims 48-55 objected to as lacking clarity under PCT Rule 66.2(a)(v) because practice of the claimed invention is not adequately described in writing, as required under PCT Rule 5.1(a)(iii), for the reasons set forth in the immediately preceding paragraph.

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.

PCT/US99/06644

Supplemental Box

(To be used when the space in any of the preceding boxes is not sufficient)

Continuation of: Boxes I - VIII

Sheet 10

CLASSIFICATION:

The International Patent Classification (IPC) and/or the National classification are as listed below:

IPC(7): A01N 63/00, A61K 39/395, C12N 15/00, A01N 61/00, C07H 21/02 and US Cl.: 424/93.1, 93.2, 130.1; 435/320.1, 325; 514/1; 536/23.1; 800/13, 18

I. BASIS OF REPORT:

5. (Some) amendments are considered to go beyond the disclosure as filed:

NONE

**MRP-Related ABC Transporter
Encoding Nucleic Acids and Methods of Use Thereof**

Pursuant to 35 U.S.C. §202(c) it is acknowledged that the U.S. Government has certain rights in the invention described herein, which was made in part with funds from the National Institutes of Health, Grant Numbers, CA63173 and CA06927.

FIELD OF THE INVENTION

The present invention relates to the fields of medicine and molecular biology. More specifically, the invention provides nucleic acid molecules and proteins encoded thereby which are involved in the development of resistance to pharmacological and chemotherapeutic agents in tumor cells.

BACKGROUND OF THE INVENTION

Several publications are referenced in this application in parentheses in order to more fully describe the state of the art to which this invention pertains. The disclosure of each of these publications is incorporated by reference herein.

P-glycoprotein, the product of the *MDR1* gene, was the first ABC transporter shown to confer resistance to cytotoxic agents. Pgp functions as an ATP-dependent efflux pump that reduces the intracellular concentration of a variety of chemotherapeutic agents by transporting them across the plasma membrane (1). The multidrug resistance phenotype associated with overexpression of Pgp

is of considerable clinical interest because natural product drugs are second only to alkylating agents in clinical utility, and many effective chemotherapeutic regimens contain more than one natural product agent. More recently, we and others have reported transfection studies indicating that MRP, another ABC family transporter, confers a multidrug resistance phenotype that includes many natural product drugs, but is distinct from the resistance phenotype associated with Pgp (2-6). MRP shares only limited amino acid identity with Pgp, and this is reflected in the different substrate specificities of the two transporters. In contrast to Pgp, MRP can transport a wide range of anionic organic conjugates, including glutathione S-conjugates (7). In addition to Pgp and MRP there may be other transporters that are involved in cytotoxic drug resistance. In the case of natural product drugs, resistant cell lines have been described that display a multidrug resistant phenotype associated with a drug accumulation deficit, but do not overexpress Pgp or MRP (8). ABC transporters have also been linked to cisplatin resistance, and several lines of evidence suggest the possibility that pumps specific for organic anions may be involved: 1) decreased cisplatin accumulation is consistently observed in cisplatin resistant cell lines (9); 2) cisplatin is conjugated to glutathione in the cell, and this anionic conjugate is toxic in an *in vitro* biochemical assay (10); and 3) biochemical studies using membrane vesicle preparations have shown that cisplatin resistant cells lines have enhanced expression of an ATP-dependent transporter of CDDP-glutathione and other glutathione S-conjugates such as the cystinyl leukotriene LTC₄ (11, 12). These data thus suggest that an organic anion transporter may contribute

to cisplatin resistance by exporting CDDP-glutathione. While MRP is an organic anion transporter, the reported drug resistance profile of MRP-transfected cells does not extend to this agent (5, 6), and to date only one cisplatin resistant cell line has been reported to overexpress MRP (13). This suggests that organic anion transporters other than MRP may contribute to cisplatin resistance. Consistent with this possibility, the canalicular multispecific organic anion transporter, cMOAT, an MRP-related transporter that functions as the major organic anion transporter in liver, has been reported to be overexpressed in cisplatin resistant cell lines (14, 15). A more direct link between cMOAT and cytotoxic drug resistance is suggested by a recent report in which transfection of a cMOAT antisense construct into a liver cancer cell line resulted in sensitization to cisplatin, daunorubicin and other cytotoxic agents (16).

Clearly, a need exists for identifying the essential components and mechanisms giving rise to drug resistance and the transport of anticancer agents out of the tumor cell. The elucidation of these mechanisms may be used to advantage for the design of efficacious chemotherapeutic agents.

SUMMARY OF THE INVENTION

This invention provides novel, biological molecules useful for identification, detection, and/or molecular characterization of components involved in the acquisition of drug resistance in tumor cells. According to one aspect of the invention, an isolated nucleic acid molecule is provided which includes a sequence encoding a protein transporter of a size between about 1300 and 1350 amino acids in length. The encoded protein, referred to herein

as MOAT-B, comprises a multi-domain structure including a tandem repeat of nucleotide binding folds appended C-terminal to a hydrophobic domain that contains several potential membrane spanning helices. Conserved Walker A and B ATP binding sites are present in each of the nucleotide binding folds.

In a preferred embodiment of the invention, an isolated nucleic acid molecule is provided that includes a cDNA encoding a human MOAT-B protein. In a particularly preferred embodiment, the human MOAT-B protein has an amino acid sequence the same as Sequence I.D. No. 2. An exemplary MOAT-B nucleic acid molecule of the invention comprises Sequence I.D. No. 1.

According to another aspect of the invention, a second isolated nucleic acid molecule is provided which includes a sequence encoding a transporter between about 1400 and 1450 amino acids. The encoded protein, referred to herein as MOAT-C contains a multi-domain structure including a tandem repeat of nucleotide binding folds appended C-terminal to a hydrophobic domain that contains several potential membrane spanning helices. Conserved Walker A and B ATP binding sites are present in each of the nucleotide binding folds. While similar in structure to MOAT-B described above, MOAT-C contains distinct sequence differences.

In a preferred embodiment of the invention, an isolated nucleic acid molecule is provided that includes a cDNA encoding a human MOAT-C protein. In a particularly preferred embodiment, the human MOAT-C protein has an amino acid sequence the same as Sequence I.D. No. 4. An exemplary MOAT-C nucleic acid molecule of the invention comprises Sequence I.D. No. 3.

According to yet another aspect of the invention, an

isolated nucleic acid molecule is provided which includes a sequence encoding a protein of a size between about 1500 and 1550 amino acids in length. The encoded protein, referred to herein as MOAT-D, contains a multidomain structure including an N-terminal hydrophobic extension which harbors five transmembrane spanning helices.

In a preferred embodiment of the invention, an isolated nucleic acid molecule is provided that includes a cDNA encoding a MOAT-D protein. In a particularly preferred embodiment, the human MOAT-D protein has an amino acid sequence the same as Sequence I.D. No. 6. An exemplary MOAT-D nucleic acid molecule of the invention comprises Sequence I.D. No. 5.

According to yet another aspect of the invention, an isolated nucleic acid molecule is provided which includes a sequence encoding a protein of a size between about 1480 and 1530 amino acids in length. The encoded protein, referred to herein as MOAT-E, contains a multidomain structure including an N-terminal hydrophobic extension which harbors several transmembrane spanning helices. While similar in structure to MOAT-D described above, MOAT-E contains distinct sequence differences.

In a preferred embodiment of the invention, an isolated nucleic acid molecule is provided that includes a cDNA encoding a MOAT-E protein. In a particularly preferred embodiment, the human MOAT-E protein has an amino acid sequence the same as Sequence I.D. No. 8. An exemplary MOAT-E nucleic acid molecule of the invention comprises Sequence I.D. No. 7.

According to another aspect of the present invention, an isolated nucleic acid molecule is provided, which has a sequence selected from the group consisting of: (1) Sequence I.D. No. 1; (2) a sequence specifically

hybridizing with preselected portions or all of the complementary strand of Sequence I.D. No. 1 comprising nucleic acids encoding amino acids 1-1154 of Sequence ID No. 2; (3) a sequence encoding preselected portions of Sequence I.D. No. 1 within nucleotides 1-3462, (4) Sequence I.D. No. 3; (5) a sequence specifically hybridizing with preselected portions or all of the complementary strand of Sequence I.D. No. 3 comprising nucleic acids encoding amino acids 1-442 of Sequence ID No. 4; (6) a sequence encoding preselected portions of Sequence I.D. No. 3 within nucleotides 1-1326, (7) Sequence I.D. No. 5; (8) a sequence specifically hybridizing with preselected portions or all of the complementary strand of Sequence I.D. No. 5 comprising nucleic acids encoding amino acids 1-1036 of Sequence ID No. 6; (9) a sequence encoding preselected portions of Sequence I.D. No. 5 within nucleotides 1-3108, (1) Sequence I.D. No. 7; (2) a sequence specifically hybridizing with preselected portions or all of the complementary strand of Sequence I.D. No. 7 comprising nucleic acids encoding amino acids 1-998 of Sequence ID No. 8; (3) a sequence encoding preselected portions of Sequence I.D. No. 7 within nucleotides 1-300.

Such partial sequences are useful as probes to identify and isolate homologues of the MOAT genes of the invention. Additionally, isolated nucleic acid sequences encoding natural allelic variants of the nucleic acids of Sequence I.D. Nos., 1, 3, 5 and 7 are also contemplated to be within the scope of the present invention. The term natural allelic variants will be defined hereinbelow.

According to another aspect of the present invention, antibodies immunologically specific for the human MOAT proteins described hereinabove are provided.

In yet another aspect of the invention, host cells comprising at least one of the MOAT encoding nucleic acids are provided. Such host cells include but are not limited to bacterial cells, fungal cells, insect cells, mammalian cells, and plant cells. Host cells overexpressing one or more of the MOAT encoding nucleic acids of the invention provide valuable research tools for assessing transport of chemotherapeutic agents out of cells. MOAT expressing cells also comprise a biological system useful in methods for identifying inhibitors of the MOAT transporters.

Another embodiment of the present invention encompasses methods for screening cells expressing MOAT encoding nucleic acids for chemotherapy resistance. Such methods will provide the clinician with data which correlates expression of a particular MOAT genes with a particular chemotherapy resistant phenotype.

Diagnostic methods are also contemplated in the present invention. Accordingly, suitable oligonucleotide probes are provided which hybridize to the nucleic acids of the invention. Such probes may be used to advantage in screening biopsy samples for the expression of particular MOAT genes. Once a tumor sample has been characterized as to the MOAT gene(s) expressed therein, inhibitors identified in the cell line screening methods described above may be administered to prevent efflux of the beneficial chemotherapeutic agents from cancer cells.

The methods of the invention may be applied to kits. An exemplary kit of the invention comprises MOAT gene specific oligonucleotide probes and/or primers, MOAT encoding DNA molecules for use as a positive control, buffers, and an instruction sheet. A kit for practicing the cell line screening method includes frozen cells

comprising the MOAT genes of the invention, suitable culture media, buffers and an instruction sheet.

In a further aspect of the invention, transgenic knockout mice are disclosed. Mice will be generated in which at least one MOAT gene has been knocked out. Such mice will provide a valuable in biological system for assessing resistance to chemotherapy in an in vivo tumor model.

Various terms relating to the biological molecules of the present invention are used hereinabove and also throughout the specification and claims. The terms "percent similarity" and "percent identity (identical)" are used as set forth in the UW GCG Sequence Analysis program (Devereux et al. NAR 12:387-397 (1984)).

With reference to nucleic acids of the invention, the term "isolated nucleic acid" is sometimes used. This term, when applied to DNA, refers to a DNA molecule that is separated from sequences with which it is immediately contiguous (in the 5' and 3' directions) in the naturally occurring genome of the organism from which it originates. For example, the "isolated nucleic acid" may comprise a DNA or cDNA molecule inserted into a vector, such as a plasmid or virus vector, or integrated into the genomic DNA of a prokaryote or eukaryote.

With respect to RNA molecules of the invention, the term "isolated nucleic acid" primarily refers to an RNA molecule encoded by an isolated DNA molecule as defined above. Alternatively, the term may refer to an RNA molecule that has been sufficiently separated from RNA molecules with which it would be associated in its natural state (i.e., in cells or tissues), such that it exists in a "substantially pure" form (the term "substantially pure" is defined below).

With respect to protein, the term "isolated protein" or "isolated and purified protein" is sometimes used herein. This term refers primarily to a protein produced by expression of an isolated nucleic acid molecule of the invention. Alternatively, this term may refer to a protein which has been sufficiently separated from other proteins with which it would naturally be associated, so as to exist in "substantially pure" form.

The term "substantially pure" refers to a preparation comprising at least 50-60% by weight the compound of interest (e.g., nucleic acid, oligonucleotide, protein, etc.). More preferably, the preparation comprises at least 75% by weight, and most preferably 90-99% by weight, the compound of interest. Purity is measured by methods appropriate for the compound of interest (e.g. chromatographic methods, agarose or polyacrylamide gel electrophoresis, HPLC analysis, and the like). With respect to antibodies of the invention, the term "immunologically specific" refers to antibodies that bind to one or more epitopes of a protein of interest (e.g., MOAT-B, MOAT-C or MOAT-D), but which do not substantially recognize and bind other molecules in a sample containing a mixed population of antigenic biological molecules.

With respect to nucleic acids and oligonucleotides, the term "specifically hybridizing" refers to the association between two single-stranded nucleotide molecules of sufficiently complementary sequence to permit such hybridization under pre-determined conditions generally used in the art (sometimes termed "substantially complementary"). When used in reference to a double stranded nucleic acid, this term is intended to signify that the double stranded nucleic acid has been subjected to denaturing conditions, as is well known to those of

skill in the art. In particular, the term refers to hybridization of an oligonucleotide with a substantially complementary sequence contained within a single-stranded DNA or RNA molecule of the invention, to the substantial exclusion of hybridization of the oligonucleotide with single-stranded nucleic acids of non-complementary sequence.

One common formula for calculating the stringency conditions required to achieve hybridization between nucleic acid molecules of a specified sequence homology (Sambrook et al., 1989):

$$T_m = 81.5^{\circ}\text{C} + 16.6\text{Log} [\text{Na}^+] + 0.41(\% \text{ G+C}) - 0.63 (\% \text{ formamide}) - 600/\text{\#bp in duplex}$$

As an illustration of the above formula, using $[\text{Na}^+] = [0.368]$ and 50% formamide, with GC content of 42% and an average probe size of 200 bases, the T_m is 57°C . The T_m of a DNA duplex decreases by $1 - 1.5^{\circ}\text{C}$ with every 1% decrease in homology. Thus, targets with greater than about 75% sequence identity would be observed using a hybridization temperature of 42°C . Such sequences would be considered substantially homologous to the nucleic acid sequences of the invention.

The nucleic acids, proteins, antibodies, cell lines, methods, and kits of the present invention may be used to advantage to identify targets for the development of novel agents which inhibit the aberrant transport of cytotoxic agents out of tumor cells. The transgenic mice of the invention may be used as an in vivo model for chemotherapy resistance.

The human MOAT molecules methods and kits described above may also be used as research tools and will facilitate the elucidation of the mechanism by which tumor

cells acquire a drug resistant phenotype.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows the predicted structure of MOAT-B and comparison with human MRP. The vertical lines indicate identical amino acids and the vertical dots indicate conserved amino acids. Gaps are indicated by periods. The overbars indicate potential transmembrane spanning segments as predicted by the TMAP program. The first and second nucleotide binding folds (NBF 1 and NBF 2) are indicated by horizontal arrows. The C-terminal 34 amino acids (residues 1291 - 1325) are replaced in the second class of MOAT-B cDNA clones by the following amino acids: ILQKKLSTYWSH. The Alignment was performed using the GAP program (gap weight 3.0, length weight 0.1) in the Genetics Computer Group Package. H. MRP: human MRP.

Figures 2A and 2B depict a comparison of the nucleotide binding folds and hydropathy profile of MOAT-B with those of other eukaryotic ABC transporters. Fig. 1A shows the comparison of the nucleotide binding folds of MOAT-B. Amino acids that are identical to those of MOAT-B are shaded, and gaps are indicated by periods. Walker A and B motifs, and the ABC transporter family signature sequence C, are underlined. Amino acid positions are indicated to the right. Amino acid sequences were aligned using the PILEUP program (gap weight 3.0, length weight 0.1) in the Genetics Computer Group Package. Fig. 1B shows a comparison of the MOAT-B hydropathy profile. To facilitate comparison, the proteins are aligned so that the N-terminal nucleotide binding folds (NBF) are roughly in register. NBF's are indicated by bars. Values above

and below the horizontal lines indicate hydrophobic and hydrophilic regions, respectively. Hydrophobicity plots were generated using the Kyte-Doolittle algorithm with a window of 7 residues. The transporters shown are: human multidrug-associated protein, H. MRP (P33529); human multispecific organic anion transporter, H. MOAT (U63970); *Saccharomyces cerevisiae* yeast cadmium factor 1, S. YCF1 (P39109); rat sulfonylurea receptor, R. SUR (Q09427); human cystic fibrosis transmembrane conductance regulator, H. CFTR (M28668); *Leishmania* P-glycoprotein, L. PgpA (P21441) and human *mdr1* gene product, H. MDR1 (P08183). Accession numbers are shown in parentheses.

Figure 3 is a Northern blot showing the tissue distribution of MOAT-B transcript. Membranes containing poly (A)+ RNA prepared from human tissues were hybridized with a radiolabeled MOAT-B or GAPDH probe. Top panels show MOAT-B transcript and bottom panels show the control GAPDH transcript. Arrows indicate the position of MOAT-B transcript. Prolonged exposure of the film revealed a low level signal in liver.

Figure 4 shows the chromosomal localization of the gene encoding MOAT-B. Human metaphase spreads were hybridized with a biotin-labeled MOAT-B cDNA probe and detected by FITC-conjugated avidin. Hybridization signals at chromosome 13q32 in two metaphase spreads are indicated by arrows. The inset shows paired hybridization signals at band q32 of chromosome 13 from three other metaphase spreads.

Figures 5A and 5B show the predicted structures of MOAT-C and MOAT-D. Fig. 5A presents the structure of

MOAT-C. Fig. 5B shows the structure of MOAT-D. Numbered overbars indicate potential transmembrane spanning helices. Horizontal arrows indicate the positions of the amino terminal (NBF1) and C-terminal (NBF2) nucleotide binding folds. Walker A and B motifs, and the ABC transporter family signature sequence C are underlined. Bullets indicate the positions of potential N-linked glycosylation sites that are conserved with previously reported N-glycosylation sites in MRP. The indicated MOAT-C transmembrane spanning helices were predicted using the TMAP program and an input alignment of MOAT-B and MOAT-C. The indicated MOAT-D transmembrane helices are based upon inspection of an alignment with MRP.

Figures 6A and 6B show a comparison of the nucleotide binding folds and hydropathy profiles of MOAT-C and MOAT-D with those of other related ABC transporters. Fig. 6A depicts the comparison of the nucleotide binding folds. The alignment was produced using the PILEUP command (gap weight 3.0, length weight 0.1) in the Genetics Computer Group Package Version 9.1. Amino acid positions conserved in at least 4 of the 8 proteins are shaded. Periods indicate gaps in the alignment. Walker A and B, and the ABC transporter family signature sequence C are indicated by underbars. Fig. 6A shows the comparison of hydropathy profiles. To facilitate comparisons, gaps were introduced at the N-termini of some proteins in order to bring the first nucleotide binding folds into register. Nucleotide binding folds are indicated by bars. Values above and below the horizontal lines indicate hydrophobic and hydrophilic regions, respectively. Hydrophobicity plots were generated using the Kyte-Doolittle algorithm with a window of 7 residues. Accession numbers are as follows:

MRP, P33529; cMOAT, U63970, SUR, Q09428; CFTR, P-13569; MDR1, P08183.

Figure 7 is a Northern blot showing the tissue distribution of MOAT-C and MOAT-D transcripts. Blots containing poly A+ RNA prepared from various human tissues were hybridized with MOAT-C, MOAT-D and actin probes. Arrows indicate the position of the MOAT-C (top panel) and MOAT-D (middle panel) transcripts. The bottom panel shows the control actin transcript.

Figures 8A and 8B show the chromosomal localization of the MOAT-C and MOAT-D genes. Human metaphase spreads were hybridized with a biotin-labeled MOAT-C and MOAT-D cDNA probes and detected by FITC-conjugated avidin. Fig. 8A shows the localization of MOAT-C. Hybridization signals at chromosome 3q27 in two metaphase spreads are indicated by arrows (top). The inset shows paired hybridization signals at band q27 of chromosome 3 from three other metaphase spreads. Fig. 8B shows the localization of MOAT-D. Hybridization signals at chromosome 17q21-22 in two metaphase spreads are indicated by arrows (top). The inset shows paired hybridization signals at band q21-22 of chromosome 17 from three other metaphase spreads.

Figure 9 shows predicted amino acid sequence of MOAT-E. Also shown are the location of the potential transmembrane helices (overbars), the potential N-glycosylation site (black dot) and the two nucleotide binding folds (NBF1 and NBF2). Walker A and B motifs, as well as the signature C motif of ABC transporters, are also indicated.

Figure 10 shows a comparison of the hydropathy profile of MOAT-E with other members of the MRP-cMOAT subfamily. The profile reveals that MOAT-E has a hydrophobic N-terminal segment which is absent in MOAT-B and MOAT-C.

Figure 11 is a RNA blot which reveals that MOAT-E is expressed only in the liver and the kidney, suggesting that MOAT-E may participate in the excretion of substances into urine and bile. The lower panel shows hybridization of an actin probe to assess RNA loading.

Figures 12A-12J show the cDNA (SEQ ID NO: 1) and amino acid sequences (SEQ ID NO: 2) encoded by MOATB.

Figures 13A-13K show the cDNA (SEQ ID NO: 3) and amino acid sequences (SEQ ID NO: 4) encoded by MOATC.

Figures 14A-14K show the cDNA (SEQ ID NO: 5) and amino acid sequences (SEQ ID NO: 6) encoded by MOATD.

Figures 15A-15K show the cDNA (SEQ ID NO: 7) and amino acid sequences (SEQ ID NO: 8) encoded by MOATE.

DETAILED DESCRIPTION OF THE INVENTION

MRP and cMOAT are closely related mammalian ABC transporters that export organic anions from cells. Transfection studies have established that MRP confers resistance to natural product cytotoxic agents, and recent evidence suggests the possibility that cMOAT may contribute to cytotoxic drug resistance as well. Based upon the potential importance of these transporters in

clinical drug resistance, and their important physiological roles in the export of the amphiphilic products of phase I and phase II metabolism, we sought to identify other MRP-related transporters. Using a degenerate PCR approach, a cDNA molecule was isolated which encodes a novel ABC transporter designated herein as MOAT-B. The MOAT-B gene was mapped using fluorescence *in situ* hybridization to chromosome band 13q32. Comparison of the MOAT-B predicted protein with other transporters revealed that it is most closely related to MRP, cMOAT, and the yeast organic anion transporter YCF1. While MOAT-B is closely related to these transporters, it is distinguished by the absence of approximately 200 amino acid N-terminal hydrophobic extension that is present in MRP and cMOAT, and which is predicted to encode several transmembrane spanning segments. In addition, the MOAT-B tissue distribution is distinct from MRP and cMOAT. In contrast to MRP, which is widely expressed in most tissues, including liver, and cMOAT, whose expression is largely restricted to liver, the MOAT-B transcript is widely expressed, with particularly high levels in prostate, but is barely detectable in liver. These data indicate that MOAT-B is a ubiquitously expressed transporter that is closely related to MRP and cMOAT, and indicate that it is an organic anion pump relevant to cellular detoxification.

Three additional MRP/cMOAT-related transporters, MOAT-C, MOAT-D and MOAT-E are also disclosed herein. MOAT-C encodes a 1437 amino acid protein that is most closely related to MRP, cMOAT and MOAT-B, among eukaryotic transporters (33% - 37% identity). However, based upon amino acid identity, MOAT-C is considerably less related to MRP and cMOAT than the latter transporters are to each

other (48% identity). In addition, the MOAT-C topology is distinct from that of MRP and cMOAT in that it, like MOAT-B, lacks an N-terminal transmembrane spanning domain. MOAT-D encodes a 1530 amino acid transporter that is highly related to MRP (57% identity) and cMOAT (47% identity). MOAT-E encodes 1503 amino acid transporter that is highly related to MOAT-D, MRP and cMOAT (39-45% identity). The topology of MOAT-D and MOAT-E are quite similar to MRP and cMOAT, in that they have an N-terminal hydrophobic extension that is predicted to harbor five transmembrane spanning helices. MOAT-C and MOAT-D were mapped to chromosome bands 3q27 and 17q21-22, respectively, by fluorescence *in situ* hybridization.

The expression patterns of MOAT-C, MOAT-D and MOAT-E are distinct from those of MRP, cMOAT and MOAT-B. MOAT-C transcript is widely expressed, with highest levels in skeletal muscle, kidney and testis, but is expressed at barely detectable levels in liver and lung. MOAT-D transcript has a more restricted expression pattern, with high levels in colon, pancreas, liver and kidney. Data presented herein reveal that MOAT-E expression is restricted to liver and kidney.

Based upon degree of amino acid identity, and protein topology, the MRP-related transporters fall into two groups, with the first group consisting of MRP, cMOAT, MOAT-D and MOAT-E, and the second group consisting of MOAT-B and MOAT-C. The isolation of MOAT-C, MOAT-D and MOAT-E thus helps to define the MRP/cMOAT subfamily. The high degree of amino acid identity and topological similarity of MOAT-D and MOAT-E to MRP and cMOAT suggest that they function as organic anion transporters, and play a role in cytotoxic drug resistance. In contrast, the lower degree of amino acid identity and distinct topology

of MOAT-B and MOAT-C suggest the possibility that their substrate specificities and functions may be distinct from that of MRP, cMOAT, MOAT-D and MOAT-E.

The compositions, methods, kits and transgenic mice of the invention disclosed herein will facilitate the identification of drugs that cripple the ability of MOAT genes and proteins encoded thereby to effect the efflux of clinically beneficial pharmacological agents in malignant cells.

I. Preparation of MOAT-Encoding Nucleic Acid Molecules, MOAT Proteins, and Antibodies Thereto

A. Nucleic Acid Molecules

Nucleic acid molecules encoding the MOAT proteins of the invention may be prepared by two general methods: (1) synthesis from appropriate nucleotide triphosphates, or (2) isolation from biological sources. Both methods utilize protocols well known in the art. The availability of nucleotide sequence information, such as cDNAs having Sequence I.D. Nos. 1, 3, 5, or 7 enables preparation of an isolated nucleic acid molecule of the invention by oligonucleotide synthesis. Synthetic oligonucleotides may be prepared by the phosphoramidite method employed in the Applied Biosystems 38A DNA Synthesizer or similar devices. The resultant construct may be purified according to methods known in the art, such as high performance liquid chromatography (HPLC). Long, double-stranded polynucleotides, such as a DNA molecule of the present invention, must be synthesized in stages, due to the size limitations inherent in current oligonucleotide synthetic methods. Thus, for example, a 5 kb double-stranded molecule may be synthesized as several smaller segments of appropriate complementarity. Complementary segments thus

produced may be annealed such that each segment possesses appropriate cohesive termini for attachment of an adjacent segment. Adjacent segments may be ligated by annealing cohesive termini in the presence of DNA ligase to construct an entire 5 kb double-stranded molecule. A synthetic DNA molecule so constructed may then be cloned and amplified in an appropriate vector.

Nucleic acid sequences encoding the MOAT proteins of the invention may be isolated from appropriate biological sources using methods known in the art. In a preferred embodiment, a cDNA clone is isolated from a cDNA expression library of human origin. In an alternative embodiment, utilizing the sequence information provided by the cDNA sequence, human genomic clones encoding MOAT proteins may be isolated. Alternatively, cDNA or genomic clones having homology with MOAT-B, MOAT-C, MOAT-D or MOAT-E may be isolated from other species using oligonucleotide probes corresponding to predetermined sequences within the MOAT encoding nucleic acids.

In accordance with the present invention, nucleic acids having the appropriate level of sequence homology with the protein coding region of Sequence I.D. Nos. 1, 3, 5, and 7 may be identified by using hybridization and washing conditions of appropriate stringency. For example, hybridizations may be performed, according to the method of Sambrook et al., (supra) using a hybridization solution comprising: 5X SSC, 5X Denhardt's reagent, 1.0% SDS, 100 µg/ml denatured, fragmented salmon sperm DNA, 0.05% sodium pyrophosphate and up to 50% formamide. Hybridization is carried out at 37-42°C for at least six hours. Following hybridization, filters are washed as follows: (1) 5 minutes at room temperature in 2X SSC and 1% SDS; (2) 15 minutes at room temperature in 2X SSC and

0.1% SDS; (3) 30 minutes-1 hour at 37°C in 1X SSC and 1% SDS; (4) 2 hours at 42-65° in 1X SSC and 1% SDS, changing the solution every 30 minutes.

Nucleic acids of the present invention may be maintained as DNA in any convenient cloning vector. In a preferred embodiment, clones are maintained in a plasmid cloning/expression vector, such as pBluescript (Stratagene, La Jolla, CA), which is propagated in a suitable *E. coli* host cell.

MOAT-encoding nucleic acid molecules of the invention include cDNA, genomic DNA, RNA, and fragments thereof which may be single- or double-stranded. Thus, this invention provides oligonucleotides (sense or antisense strands of DNA or RNA) having sequences capable of hybridizing with at least one sequence of a nucleic acid molecule of the present invention, such as selected segments of the cDNA having Sequence I.D. No. 1. Such oligonucleotides are useful as probes for detecting or isolating MOAT genes. Antisense nucleic acid molecules may be targeted to translation initiation sites and/or splice sites to inhibit the translation of the MOAT-encoding nucleic acids of the invention. Such antisense molecules are typically between 15 and 30 nucleotides in length and often span the translational start site of MOAT encoding mRNA molecules.

It will be appreciated by persons skilled in the art that variants of these sequences exist in the human population, and must be taken into account when designing and/or utilizing oligos of the invention. Accordingly, it is within the scope of the present invention to encompass such variants, with respect to the MOAT sequences disclosed herein or the oligos targeted to specific locations on the respective genes or RNA transcripts.

With respect to the inclusion of such variants, the term "natural allelic variants" is used herein to refer to various specific nucleotide sequences and variants thereof that would occur in a human population. The usage of different wobble codons and genetic polymorphisms which give rise to conservative or neutral amino acid substitutions in the encoded protein are examples of such variants. Additionally, the term "substantially complementary" refers to oligo sequences that may not be perfectly matched to a target sequence, but the mismatches do not materially affect the ability of the oligo to hybridize with its target sequence under the conditions described.

B. Proteins

Full-length MOAT-B, MOAT-C, MOAT-D and MOAT-E proteins of the present invention may be prepared in a variety of ways, according to known methods. The proteins may be purified from appropriate sources, e.g., transformed bacterial or animal cultured cells or tissues, by immunoaffinity purification. However, this is not a preferred method due to the low amount of protein likely to be present in a given cell type at any time. The availability of nucleic acid molecules encoding MOAT proteins enables production of the proteins using *in vitro* expression methods known in the art. For example, a cDNA or gene may be cloned into an appropriate *in vitro* transcription vector, such as pSP64 or pSP65 for *in vitro* transcription, followed by cell-free translation in a suitable cell-free translation system, such as wheat germ or rabbit reticulocytes. *In vitro* transcription and translation systems are commercially available, e.g., from Promega Biotech, Madison, Wisconsin or Gibco-BRL,

Gaithersburg, Maryland.

Alternatively, according to a preferred embodiment, larger quantities of MOAT proteins may be produced by expression in a suitable prokaryotic or eukaryotic system. For example, part or all of a DNA molecule, such as a cDNA having Sequence I.D. No. 1, 3, 5 or 7 may be inserted into a plasmid vector adapted for expression in a bacterial cell, such as *E. coli*. Such vectors comprise the regulatory elements necessary for expression of the DNA in the host cell positioned in such a manner as to permit expression of the DNA in the host cell. Such regulatory elements required for expression include promoter sequences, transcription initiation sequences and, optionally, enhancer sequences.

The human MOAT proteins produced by gene expression in a recombinant procaryotic or eukaryotic system may be purified according to methods known in the art. In a preferred embodiment, a commercially available expression/secretion system can be used, whereby the recombinant protein is expressed and thereafter secreted from the host cell, to be easily purified from the surrounding medium. If expression/secretion vectors are not used, an alternative approach involves purifying the recombinant protein by affinity separation, such as by immunological interaction with antibodies that bind specifically to the recombinant protein or nickel columns for isolation of recombinant proteins tagged with 6-8 histidine residues at their N-terminus or C-terminus. Alternative tags may comprise the FLAG epitope or the hemagglutinin epitope. Such methods are commonly used by skilled practitioners.

The human MOAT proteins of the invention, prepared by the aforementioned methods, may be analyzed according to

standard procedures. For example, such proteins may be subjected to amino acid sequence analysis, according to known methods.

The present invention also provides antibodies capable of immunospecifically binding to proteins of the invention. Polyclonal antibodies directed toward human MOAT proteins may be prepared according to standard methods. In a preferred embodiment, monoclonal antibodies are prepared, which react immunospecifically with the various epitopes of the MOAT proteins described herein. Monoclonal antibodies may be prepared according to general methods of Köhler and Milstein, following standard protocols. Polyclonal or monoclonal antibodies that immunospecifically interact with MOAT proteins can be utilized for identifying and purifying such proteins. For example, antibodies may be utilized for affinity separation of proteins with which they immunospecifically interact. Antibodies may also be used to immunoprecipitate proteins from a sample containing a mixture of proteins and other biological molecules. Other uses of anti-MOAT antibodies are described below.

II. Uses of MOAT-Encoding Nucleic Acids, MOAT Proteins and Antibodies Thereto

Cellular transporter molecules have received a great deal of attention as potential targets of chemotherapeutic agents designed to effectively block the export of pharmacological reagents from tumor cells. The MOAT proteins of the invention play a pivotal role in the transport of molecules across the cell membrane.

Additionally, MOAT nucleic acids, proteins and antibodies thereto, according to this invention, may be used as research tools to identify other proteins that are

intimately involved in the transport of molecules into and out of cells. Biochemical elucidation of molecular mechanisms which govern such transport will facilitate the development of novel anti-transport agents that may sensitize tumor cells to conventional chemotherapeutic agents.

A. MOAT-Encoding Nucleic Acids

MOAT-encoding nucleic acids may be used for a variety of purposes in accordance with the present invention. MOAT-encoding DNA, RNA, or fragments thereof may be used as probes to detect the presence of and/or expression of genes encoding MOAT proteins. Methods in which MOAT-encoding nucleic acids may be utilized as probes for such assays include, but are not limited to: (1) *in situ* hybridization; (2) Southern hybridization (3) northern hybridization; and (4) assorted amplification reactions such as polymerase chain reactions (PCR).

The MOAT-encoding nucleic acids of the invention may also be utilized as probes to identify related genes from other animal species. As is well known in the art, hybridization stringencies may be adjusted to allow hybridization of nucleic acid probes with complementary sequences of varying degrees of homology. Thus, MOAT-encoding nucleic acids may be used to advantage to identify and characterize other genes of varying degrees of relation to the MOAT genes of the invention. Such information enables further characterization of transporter molecules which give rise to the chemoresistant phenotype of certain tumors. Additionally, they may be used to identify genes encoding proteins that interact with MOAT proteins (e.g., by the "interaction trap" technique), which should further accelerate

identification of the components involved in the acquisition of drug resistance. The MOAT encoding nucleic acids may also be used to generate primer sets suitable for PCR amplification of target MOAT DNA. Criteria for selecting suitable primers are well known to those of ordinary skill in the art.

Nucleic acid molecules, or fragments thereof, encoding MOAT genes may also be utilized to control the production of MOAT proteins, thereby regulating the amount of protein available to participate in cytotoxic drug efflux. As mentioned above, antisense oligonucleotides corresponding to essential processing sites in MOAT-encoding mRNA molecules may be utilized to inhibit MOAT protein production in targeted cells. Alterations in the physiological amount of MOAT proteins may dramatically affect the ability of these proteins to transport pharmacological reagents out of the cell.

Host cells comprising at least one MOAT encoding DNA molecule are encompassed in the present invention. Host cells contemplated for use in the present invention include but are not limited to bacterial cells, fungal cells, insect cells, mammalian cells, and plant cells. The MOAT encoding DNA molecules may be introduced singly into such host cells or in combination to assess the phenotype of cells conferred by such expression. Methods for introducing DNA molecules are also well known to those of ordinary skill in the art. Such methods are set forth in Ausubel et al. eds., Current Protocols in Molecular Biology, John Wiley & Sons, NY, NY 1995, the disclosure of which is incorporated by reference herein.

The availability of MOAT encoding nucleic acids enables the production of strains of laboratory mice carrying part or all of the MOAT genes or mutated

sequences thereof. Such mice may provide an in vivo model for development of novel chemotherapeutic agents.

Alternatively, the MOAT nucleic acid sequence information provided herein enables the production of knockout mice in which the endogenous genes encoding MOAT-B, MOAT-C, MOAT-D or MOAT-E have been specifically inactivated. Methods of introducing transgenes in laboratory mice are known to those of skill in the art. Three common methods include: 1. integration of retroviral vectors encoding the foreign gene of interest into an early embryo; 2. injection of DNA into the pronucleus of a newly fertilized egg; and 3. the incorporation of genetically manipulated embryonic stem cells into an early embryo.

The alterations to the MOAT gene envisioned herein include modifications, deletions, and substitutions. Modifications and deletions render the naturally occurring gene nonfunctional, producing a "knock out" animal. Substitutions of the naturally occurring gene for a gene from a second species results in an animal which produces an MOAT gene from the second species. Substitution of the naturally occurring gene for a gene having a mutation results in an animal with a mutated MOAT protein. A transgenic mouse carrying the human MOAT gene is generated by direct replacement of the mouse MOAT gene with the human gene. These transgenic animals are valuable for use in vivo assays for elucidation of other medical disorders associated with cellular activities modulated by MOAT genes. A transgenic animal carrying a "knock out" of a MOAT encoding nucleic acid is useful for the establishment of a nonhuman model for chemotherapy resistance involving MOAT regulation.

As a means to define the role that MOAT plays in mammalian systems, mice can be generated that cannot make

MOAT proteins because of a targeted mutational disruption of a MOAT gene.

The term "animal" is used herein to include all vertebrate animals, except humans. It also includes an individual animal in all stages of development, including embryonic and fetal stages. A "transgenic animal" is any animal containing one or more cells bearing genetic information altered or received, directly or indirectly, by deliberate genetic manipulation at the subcellular level, such as by targeted recombination or microinjection or infection with recombinant virus. The term "transgenic animal" is not meant to encompass classical cross-breeding or in vitro fertilization, but rather is meant to encompass animals in which one or more cells are altered by or receive a recombinant DNA molecule. This molecule may be specifically targeted to defined genetic locus, be randomly integrated within a chromosome, or it may be extrachromosomally replicating DNA. The term "germ cell line transgenic animal" refers to a transgenic animal in which the genetic alteration or genetic information was introduced into a germ line cell, thereby conferring the ability to transfer the genetic information to offspring. If such offspring in fact, possess some or all of that alteration or genetic information, then they, too, are transgenic animals.

The alteration or genetic information may be foreign to the species of animal to which the recipient belongs, or foreign only to the particular individual recipient, or may be genetic information already possessed by the recipient. In the last case, the altered or introduced gene may be expressed differently than the native gene.

The altered MOAT gene generally should not fully encode the same MOAT protein native to the host animal and

its expression product should be altered to a minor or great degree, or absent altogether. However, it is conceivable that a more modestly modified MOAT gene will fall within the compass of the present invention if it is a specific alteration.

The DNA used for altering a target gene may be obtained by a wide variety of techniques that include, but are not limited to, isolation from genomic sources, preparation of cDNAs from isolated mRNA templates, direct synthesis, or a combination thereof.

A preferred type of target cell for transgene introduction is the embryonal stem cell (ES). ES cells may be obtained from pre-implantation embryos cultured in vitro. Transgenes can be efficiently introduced into the ES cells by standard techniques such as DNA transfection or by retrovirus-mediated transduction. The resultant transformed ES cells can thereafter be combined with blastocysts from a non-human animal. The introduced ES cells thereafter colonize the embryo and contribute to the germ line of the resulting chimeric animal.

One approach to the problem of determining the contributions of individual genes and their expression products is to use isolated MOAT genes to selectively inactivate the wild-type gene in totipotent ES cells (such as those described above) and then generate transgenic mice. The use of gene-targeted ES cells in the generation of gene-targeted transgenic mice is known in the art.

Techniques are available to inactivate or alter any genetic region to a mutation desired by using targeted homologous recombination to insert specific changes into chromosomal alleles. However, in comparison with homologous extrachromosomal recombination, which occurs at a frequency approaching 100%, homologous plasmid-

chromosome recombination was originally reported to only be detected at frequencies between 10^{-6} and 10^{-3} .

Nonhomologous plasmid-chromosome interactions are more frequent occurring at levels 10^5 -fold to 10^2 -fold greater than comparable homologous insertion.

To overcome this low proportion of targeted recombination in murine ES cells, various strategies have been developed to detect or select rare homologous recombinants. One approach for detecting homologous alteration events uses the polymerase chain reaction (PCR) to screen pools of transformant cells for homologous insertion, followed by screening of individual clones. Alternatively, a positive genetic selection approach has been developed in which a marker gene is constructed which will only be active if homologous insertion occurs, allowing these recombinants to be selected directly. One of the most powerful approaches developed for selecting homologous recombinants is the positive-negative selection (PNS) method developed for genes for which no direct selection of the alteration exists. The PNS method is more efficient for targeting genes which are not expressed at high levels because the marker gene has its own promoter. Non-homologous recombinants are selected against by using the Herpes Simplex virus thymidine kinase (HSV-TK) gene and selecting against its nonhomologous insertion with effective herpes drugs such as gancyclovir (GANC) or (1-(2-deoxy-2-fluoro-B-D arabinofluranosyl)-5-iodouracil, (FIAU). By this counter selection, the number of homologous recombinants in the surviving transformants can be increased.

As used herein, a "targeted gene" or "knock-out" is a DNA sequence introduced into the germline or a non-human animal by way of human intervention, including but not

limited to, the methods described herein. The targeted genes of the invention include DNA sequences which are designed to specifically alter cognate endogenous alleles.

Methods of use for the transgenic mice of the invention are also provided herein. Knockout mice of the invention can be injected with tumor cells or treated with carcinogens to generate carcinomas. Such mice provide a biological system for assessing chemotherapy resistance as modulated by a MOAT gene of the invention. Accordingly, therapeutic agents which inhibit the action of these transporters and thereby prevent efflux of beneficial chemotherapeutic agents from tumor cells may be screened in studies using MOAT knock out mice.

As described above, MOAT-encoding nucleic acids are also used to advantage to produce large quantities of substantially pure MOAT proteins, or selected portions thereof.

B. MOAT Proteins and Antibodies

Purified full length MOAT proteins, or fragments thereof, may be used to produce polyclonal or monoclonal antibodies which also may serve as sensitive detection reagents for the presence and accumulation of MOAT proteins (or complexes containing MOAT proteins) in mammalian cells. Recombinant techniques enable expression of fusion proteins containing part or all of MOAT proteins. The full length proteins or fragments of the proteins may be used to advantage to generate an array of monoclonal antibodies specific for various epitopes of MOAT proteins, thereby providing even greater sensitivity for detection of MOAT proteins in cells.

Polyclonal or monoclonal antibodies immunologically specific for MOAT proteins may be used in

a variety of assays designed to detect and quantitate the proteins. Such assays include, but are not limited to:

(1) flow cytometric analysis; (2) immunochemical localization of MOAT proteins in tumor cells; and (3) immunoblot analysis (e.g., dot blot, Western blot) of extracts from various cells. Additionally, as described above, anti-MOAT antibodies can be used for purification of MOAT proteins and any associated subunits (e.g., affinity column purification, immunoprecipitation).

From the foregoing discussion, it can be seen that MOAT-encoding nucleic acids, MOAT expressing vectors, MOAT proteins and anti-MOAT antibodies of the invention can be used to detect MOAT gene expression and alter MOAT protein accumulation for purposes of assessing the genetic and protein interactions involved in the development of drug resistance in tumor cells.

C. Methods and Kits Employing the

Compositions of the Present Invention

From the foregoing discussion, it can be seen that MOAT-encoding nucleic acids, MOAT-expressing vectors, MOAT proteins and anti-MOAT antibodies of the invention can be used to detect MOAT gene expression and alter MOAT protein accumulation for purposes of assessing the genetic and protein interactions giving rise to chemotherapy resistance in tumor cells.

Exemplary approaches for detecting MOAT nucleic acid or polypeptides/proteins include:

- a) comparing the sequence of nucleic acid in the sample with the MOAT nucleic acid sequence to determine whether the sample from the patient contains mutations; or
- b) determining the presence, in a sample from a patient, of the polypeptide encoded by the MOAT gene and,

if present, determining whether the polypeptide is full length, and/or is mutated, and/or is expressed at the normal level; or

c) using DNA restriction mapping to compare the restriction pattern produced when a restriction enzyme cuts a sample of nucleic acid from the patient with the restriction pattern obtained from normal MOAT gene or from known mutations thereof; or,

d) using a specific binding member capable of binding to a MOAT nucleic acid sequence (either normal sequence or known mutated sequence), the specific binding member comprising nucleic acid hybridizable with the MOAT sequence, or substances comprising an antibody domain with specificity for a native or mutated MOAT nucleic acid sequence or the polypeptide encoded by it, the specific binding member being labelled so that binding of the specific binding member to its binding partner is detectable; or,

e) using PCR involving one or more primers based on normal or mutated MOAT gene sequence to screen for normal or mutant MOAT gene in a sample from a patient.

A "specific binding pair" comprises a specific binding member (sbm) and a binding partner (bp) which have a particular specificity for each other and which in normal conditions bind to each other in preference to other molecules. Examples of specific binding pairs are antigens and antibodies, ligands and receptors and complementary nucleotide sequences. The skilled person is aware of many other examples and they do not need to be listed here. Further, the term "specific binding pair" is also applicable where either or both of the specific binding member and the binding partner comprise a part of a large molecule. In embodiments in which the specific

binding pair are nucleic acid sequences, they will be of a length to hybridize to each other under conditions of the assay, preferably greater than 10 nucleotides long, more preferably greater than 15 or 20 nucleotides long.

In most embodiments for screening for alleles giving rise to chemotherapy resistance, the MOAT nucleic acid in biological sample will initially be amplified, e.g. using PCR, to increase the amount of the analyte as compared to other sequences present in the sample. This allows the target sequences to be detected with a high degree of sensitivity if they are present in the sample. This initial step may be avoided by using highly sensitive array techniques that are becoming increasingly important in the art.

The identification of the MOAT gene and its association with a particular chemotherapy resistance paves the way for aspects of the present invention to provide the use of materials and methods, such as are disclosed and discussed above, for establishing the presence or absence in a test sample of a variant form of the gene, in particular an allele or variant specifically associated with chemotherapy resistance. This may be done to assess the propensity of the tumor to exhibit chemotherapy resistance.

In still further embodiments, the present invention concerns immunodetection methods for binding, purifying, removing, quantifying or otherwise generally detecting biological components. The encoded proteins or peptides of the present invention may be employed to detect antibodies having reactivity therewith, or, alternatively, antibodies prepared in accordance with the present invention, may be employed to detect the encoded proteins or peptides. The steps of various useful immunodetection methods have been

described in the scientific literature, such as, e.g., Nakamura et al. (1987).

In general, the immunobinding methods include obtaining a sample suspected of containing a protein, peptide or antibody, and contacting the sample with an antibody or protein or peptide in accordance with the present invention, as the case may be, under conditions effective to allow the formation of immunocomplexes.

The immunobinding methods include methods for detecting or quantifying the amount of a reactive component in a sample, which methods require the detection or quantitation of any immune complexes formed during the binding process. Here, one would obtain a sample suspected of containing a MOAT gene encoded protein, peptide or a corresponding antibody, and contact the sample with an antibody or encoded protein or peptide, as the case may be, and then detect or quantify the amount of immune complexes formed under the specific conditions.

In terms of antigen detection, the biological sample analyzed may be any sample that is suspected of containing the MOAT antigen, such as a tumor tissue section or specimen, a homogenized tissue extract, an isolated cell, a cell membrane preparation, separated or purified forms of any of the above protein-containing compositions.

Contacting the chosen biological sample with the protein, peptide or antibody under conditions effective and for a period of time sufficient to allow the formation of immune complexes (primary immune complexes) is generally a matter of simply adding the composition to the sample and incubating the mixture for a period of time long enough for the antibodies to form immune complexes with, i.e., to bind to, any antigens present. After this time, the sample-antibody composition, such as a tissue

section, ELISA plate, dot blot or Western blot, will generally be washed to remove any non-specifically bound antibody species, allowing only those antibodies specifically bound within the primary immune complexes to be detected.

In general, the detection of immunocomplex formation is well known in the art and may be achieved through the application of numerous approaches. These methods are generally based upon the detection of a label or marker, such as any radioactive, fluorescent, biological or enzymatic tags or labels of standard use in the art. U.S. Patents concerning the use of such labels include U.S. Pat. Nos. 3,817,837; 3,850,752; 3,939,350; 3,996,345; 4,277,437; 4,275,149 and 4,366,241, each incorporated herein by reference. Of course, one may find additional advantages through the use of a secondary binding ligand such as a second antibody or a biotin/avidin ligand binding arrangement, as is known in the art.

In one broad aspect, the present invention encompasses kits for use in detecting expression of MOAT encoding nucleic acids in biological samples, including biopsy samples. Such a kit may comprise one or more pairs of primers for amplifying nucleic acids corresponding to the MOAT gene. The kit may further comprise samples of total mRNA derived from tissues expressing at least one or a subset of the MOAT genes of the invention, to be used as controls. The kit may also comprise buffers, nucleotide bases, and other compositions to be used in hybridization and/or amplification reactions. Each solution or composition may be contained in a vial or bottle and all vials held in close confinement in a box for commercial sale. In a further embodiment, the invention encompasses a kit for use in detecting MOAT proteins in chemotherapy

resistant cancer cells comprising antibodies specific for MOAT proteins encoded by the MOAT nucleic acids of the present invention.

Another aspect of the present invention comprises screening methods employing host cells expressing one or more MOAT genes of the invention. An advantage of having discovered the complete coding sequenced of MOAT B-E is that cell lines that overexpress MOATB C D or E can be generated using standard transfection protocols. Cells that overexpress the complete cDNA will also harbor the complete proteins, a feature that is essential for biological activity of proteins. The overexpressing cell lines will be useful in several ways: 1)The drug sensitivity of overexpressing cell lines can be tested with a variety of known anticancer agents in order to determine the spectrum of anticancer agents for which the transporter confers resistance; 2)The drug sensitivity of overexpressing cell lines can be used to determine whether newly discovered anticancer agents are transported out of the cell by one of the discovered transporters; 3)Overexpressing cell lines can be used to identify potential inhibitors that reduce the activity of the transporters. Such inhibitors are of great clinical interest in that they may enhance the activity of known anticancer agents, thereby increasing their effectiveness. Reduced activity will be detected by restoration of anticancer drug sensitivity, or by reduction of transporter mediated cellular efflux of anticancer agents. In vitro biochemical studies designed to identify reduced transporter activity in the presence of potential inhibitors can also be performed using membranes prepared from overexpressing cell lines; and 4)Overexpressing cell lines can also be used to

determine whether pharmaceutical agents that are not anticancer agents are transported out of the cell by the transporters.

The following protocols are provided to facilitate the practice of the present invention.

Isolation of MOAT-B cDNA

Forward {CT(A/G/T) GT(A/G/T) GC(A/G/T) GT(A/G/T) GT(A/G/T) GG(A/G/C/T)} (SEQ ID NO:9) and reverse {(G/A)CT (A/G/C/T)A(A/G/C) (A/G/C/T)GC (A/G/C/T)(G/C)(T/A) (A/G/C/T)A(A/G) (A/G/C/T)GG (A/G/C/T)TC (A/G)TC} (SEQ ID NO:16) degenerate oligonucleotide primers were designed based upon the first nucleotide binding folds of human MRP, CFTR, and MDR1. Bacteriophage DNA isolated from a C200 cDNA library prepared in the λ pCEV27 phagemid vector (17) was used as template in PCR reactions containing 250 ng cDNA, 5 μ M primers, 50 mM KCl, 10 mM Tris-HCl, pH 8.3, 3 mM MgCl₂, .05% gelatin, 0.2 mM dNTP and Taq polymerase (Perkin Elmer Cetus). Five cycles of PCR were performed as follows: 94°C for 1 minute, 40°C for 2 minutes, 72°C for 3 minutes. Twenty five cycles were then performed as follows: 94°C for 1 minute, 55°C for 1 minute, and 72°C for 1 minute. The resulting reaction products were used as template in a second round of PCR, as described above, with nested forward {CGGGATCC AG(A/G) GA(A/G) AA(C/T) AT(A/C/T) CT(A/G/C/T) TTT GG(A/G/C/T)} (SEQ ID NO:17) and reverse {CGGAATTC (A/G/T/C)TC (A/G)TC (A/C/T)AG (A/G/C/T)AG (A/G)TA (A/T/G)AT (A/G)TC} (SEQ ID NO:18) degenerate oligonucleotide primers. PCR reaction products were isolated from an agarose gel and subcloned into the BamHI and EcoRI sites of pBluescript (Stratagene). Nucleotide sequence analysis

was performed on plasmid DNA prepared from ampicillin resistant transformants. Additional cDNA clones were isolated from C200 (ovary) and B5 (breast) cDNA libraries by plaque hybridization using the PCR product as the initial radiolabeled probe.

RNA Blot Analysis

Blots containing polyA⁺ RNA isolated from human tissues (Clontech) were prehybridized at 45°C for 8 hours in 50% formamide, 4X SSC, 4X Denhardt's solution, 0.04 M sodium phosphate monobasic, pH 6.5, 0.8% (w/v) glycine, 0.1 mg/ml sheared denatured salmon sperm DNA. Hybridization was performed at 45°C with ³²P-labeled MOAT-B or GAPDH probes in a solution containing 50% formamide, 3X SSC, 0.04 M sodium phosphate pH 6.5, 10% dextran sulfate, 0.1 mg/ml sheared denatured salmon sperm DNA. Blots were washed 2 times for 15 min at 65°C in 2X SSC, 5 mM Tris-HCl pH 7.4, 0.5% SDS, 2.5 mM EDTA, 0.1% sodium pyrophosphate pH 8.0, and subsequently washed 2 times for 15 min in 0.1X SSC. Blots were then subjected to autoradiography.

Chromosomal localization

Preparation of metaphase spreads from phytohemagglutinin-stimulated lymphocytes of a healthy female donor, and fluorescence *in situ* hybridization and detection of immunofluorescence were carried out as previously described (18). A 2.2-kb cDNA clone of MOAT-B inserted in pBluescript was biotinylated by nick translation in a reaction containing 1 µg DNA, 20 µM each of dATP, dCTP and dGTP, 1 µM dTTP, 25 mM Tris-HCl, pH 7.5, 5 mM MgCl₂, 10 mM β-mercaptoethanol, 10 µM biotin-16-dUTP (Boehringer Mannheim), 2 units DNA polymerase 1/DNase 1 (GIBCO, BRL) and water to a total volume of 50 µl. The

probe was denatured and hybridized to metaphase spreads overnight at 37°C. Hybridization sites were detected with fluorescein-labeled avidin (Oncor) and amplified by addition of anti-avidin antibody (Oncor) and a second layer of fluorescein-labeled avidin. The chromosome preparations were counterstained with DAPI and observed with a Zeiss Axiophot epifluorescence microscope equipped with a cooled charge coupled device camera (Photometrics, Tucson AZ) operated by a Macintosh computer work station. Digitized images of DAPI staining and fluorescein signals were captured, pseudo-colored and merged using Oncor Image version 1.6 software.

Isolation of MOAT-C and MOAT-D cDNA

MOAT-C and MOAT-D cDNA clones were isolated by plaque hybridization from bacteriophage cDNA libraries using the I.M.A.G.E. clones as the initial probes (ATCC).

RNA blot analysis

Blots containing polyA⁺ RNA isolated from human tissues (Clontech) were purchased from Clontech, and hybridized with radiolabeled MOAT-C, MOAT-D or actin probes according to the manufacturer's directions.

Chromosomal localization

Preparation of metaphase spreads from phytohemagglutinin-stimulated lymphocytes of a healthy female donor, and fluorescence *in situ* hybridization and detection of immunofluorescence were carried out as previously described (18). A MOAT-C probe inserted in pBluescript, or MOAT-D probe inserted in pBluescript, was biotinylated by nick translation in a reaction containing 1 µg DNA, 20 µM each of dATP, dCTP and dGTP, 1 µM dTTP, 25

mM Tris-HCl, pH 7.5, 5 mM MgCl₂, 10 mM β -mercaptoethanol, 10 μ M biotin-16-dUTP (Boehringer Mannheim), 2 units DNA polymerase I/DNase I (GIBCO, BRL) and water to a total volume of 50 μ l. The probe was denatured and hybridized to metaphase spreads overnight at 37°C. Hybridization sites were detected with fluorescein-labeled avidin (Oncor) and amplified by addition of anti-avidin antibody (Oncor) and a second layer of fluorescein-labeled avidin. The chromosome preparations were counterstained with DAPI and observed with a Zeiss Axiophot epifluorescence microscope equipped with a cooled charge coupled device camera (Photometrics, Tucson AZ) operated by a Macintosh computer work station. Digitized images of DAPI staining and fluorescein signals were captured, pseudo-colored and merged using Oncor Image version 1.6 software.

The following examples are provided to illustrate various embodiments of the invention. They are not intended to limit the invention in any way.

EXAMPLE I

Isolation of MOAT-B cDNA.

A degenerate PCR approach was used to isolate MRP-related transporters. Degenerate oligonucleotide primers were prepared based upon the N-terminal nucleotide binding folds of MRP and other eukaryotic transporters, and used in conjunction with DNA prepared from an ovarian cancer cell line bacteriophage library. Nucleotide sequence analysis of one of the resulting PCR products indicated that it encoded a segment of a novel nucleotide binding fold that was most closely related to MRP and cMOAT. Overlapping cDNA clones were isolated from ovarian and breast bacteriophage libraries by plaque hybridization using the PCR product as the initial probe. A total of

5.9 kB of cDNA was isolated. Nucleotide sequence analysis revealed two classes of cDNA clones that were about equally represented among isolates from each of the two bacteriophage libraries. The first class contained an open reading frame of 3975 bp that was bordered by in frame stop codons located at positions -76 and -42 (relative to the putative initiation codon) and 3976, and encoding a predicted protein of 1325 amino acids, which is designated MOAT-B. The open reading frame was followed by approximately 2 kB of 3' untranslated sequences. The most upstream ATG in the open reading frame was located in the sequence context 'CAAGATGC'. The A at position -3 of the putative translation initiation codon was in agreement with the major feature of the Kozak consensus sequence, but the C at position +4 was divergent from the more usual G. The second class of cDNA clones was identical to the first with the exception of a single nucleotide. These clones harbored an additional T following nucleotide 3872 of the first class of clones, close to the C-terminus of the predicted protein. This additional nucleotide resulted in a frame shift such that the predicted protein of the second class of cDNA clones was 22 residues shorter than that of the first class of cDNA clones, and in which the C-terminal 34 residues of the latter reading frame were replaced by 12 distinct residues. See brief description of Figure 1.

Analysis of the MOAT-B Predicted Structure.

Comparison of the MOAT-B predicted protein with complete coding sequences in protein data bases using the BLAST program indicated that it shared significant similarity with several eukaryotic ABC transporters. Table I.

Table I. Comparison of peptide domains of MOAT-B with those of other eukaryotic ABC transporters

MOAT-B Domain (peptide)	TM1 (88-376)	NBF1 (428-576)	linker region (577-705)	TM2 (706-992)	NBF2 (1058- 1216)	C- terminus (1217- 1325)	overall identity
percent identity							
MRP human	28.6	55.6	27.9	33.3	61.6	51.6	39.2
YCF1 yeast	27	56	27.9	34	57.2	48.5	38.9
MOAT human	33.2	53.3	32.8	31.4	55.3	44.9	38
CFTR Human	30.5	48	27.9	37.7	44	21	36.3
SUR rat	28.1	41.3	28.2	30	52.8	42.8	32.9
MDR1 human	17.6	39.2	21.1	17.3	32.2	40.3	23.3

^B The indicated domains are, TM1: segment containing the transmembrane spanning domain N-terminal to NBF1; NBF1 and NBF2: nucleotide binding folds 1 and 2; Linker region: segment located between NBF1 and TM2; TM2: segment containing the transmembrane spanning domain located between the two NBFs; C-terminus: segment between NBF2 and the C-terminus of the proteins. Sequence alignments were generated using the PILEUP program of the GCC package. Percent amino acid identity with MOAT-B domains are shown.

Typical features of eukaryotic ABC transporters were present in the predicted MOAT-B protein. See Figure 1. Overall the protein was composed of a tandem repeat of a nucleotide binding fold appended C-terminal to a hydrophobic domain that contained several potential transmembrane spanning helices. Conserved Walker A and B ATP binding sites were present in each of the nucleotide binding folds. See Figure 2A. In addition, a conserved C motif, the signature sequence of ABC transporters, was present in each nucleotide binding fold. Analysis of potential transmembrane motifs using the TMAP program (19) and an input sequence alignment of MOAT-B and MOAT-C, a transporter highly related to MOAT-B⁴, predicted 12 transmembrane helices with 6 transmembrane segments in

each of the two hydrophobic domains. This 6 + 6 configuration of predicted transmembrane helices is in agreement with topological models proposed for MRP and other ABC transporters (20, 21), and is shown in Figure 1. However, alternative predictions of transmembrane segments were obtained using different program parameters or input sequence alignments. For example, when the TMAP program was used with an input sequence alignment consisting of human MRP, rat cMOAT, rat sulfonyl urea receptor (SUR), human cystic fibrosis conductance regulator (CFTR) and human P-glycoprotein, a 6 + 5 configuration was predicted. The only substantial difference between the latter prediction and the structure shown in Figure 1 is that transmembrane segments 9 (829-853) and 10 (855-878) were replaced by a single predicted transmembrane segment spanning amino acids 847 - 875.

Among ABC transporters, the degree of similarity of the nucleotide binding folds is considered to be the best indicator of functional conservation. Comparison of the nucleotide binding folds of MOAT-B with other eukaryotic ABC transporters indicated that it was most closely related to MRP, the yeast cadmium resistance protein (YCF1) and cMOAT (Table I), three transporters that have organic anions as substrates. The MOAT-B NBF1 was 55.6, 56.0 and 53.3 percent identical, and the MOAT-B NBF2 was 61.6, 57.2 and 55.3 percent identical to the first and second nucleotide binding folds of human MRP, YCF1 and human cMOAT, respectively. Aside from the latter transporters, the MOAT-B nucleotide binding folds were most closely related to those of CFTR and SUR. The MOAT-B nucleotide binding folds shared significantly less similarity with those of MDR1. Alignment of the MOAT-B nucleotide binding folds with those of other eukaryotic

transporters is shown in Figure 2A. Analysis of the overall amino acid identity of MOAT-B with other ABC transporters also indicated that it was most closely related to MRP, YCF1 and cMOAT (Table I). Overall MOAT-B was 39.2, 38.9 and 38 percent identical to these transporters, respectively. Figure 2B shows a comparison of the hydropathy profiles of MOAT-B with those of other eukaryotic transporters. This comparison reveals that MOAT-B (1325 amino acids) is approximately 200 amino acids smaller than MRP (1531 residues), cMOAT (1545 residues) and YCF1 (1515 residues), and that this size difference is largely accounted for by the absence in MOAT-B of an amino terminal hydrophobic extension that is present in MRP, cMOAT and YCF1 (22). This N-terminal hydrophobic segment is predicted to harbor several transmembrane spanning segments, and is also present in SUR.

Expression Pattern of MOAT-B in Human Tissues.

To gain insight into the possible function of MOAT-B, its expression pattern in a variety of human tissues was examined by RNA blot analysis. As shown in Figure 3, a MOAT-B transcript of approximately 6 kB was readily detected. The isolation of 5.9 kB of MOAT-B cDNA was consistent with this size. MOAT-B expression was detected in each of the 16 tissues analyzed. Transcript levels were highest in prostate and lowest in liver and peripheral blood leukocytes, for which prolonged exposure of film were required to detect expression. Intermediate levels of expression were observed in other tissues.

Chromosomal Localization of the MOAT-B Gene.

The MOAT-B chromosomal localization was determined by fluorescence *in situ* hybridization. As shown in Figure 4, hybridization of the MOAT-B probe to metaphase spreads revealed specific labeling at human chromosome band 13q32.

Fluorescent signals were detected on chromosome 13 in each of 19 metaphase spreads scored. Of 135 signals observed, 62 (46%) were on 13q. Among these signals, 61 localized at 13q32, near the boundary between 13q31 and 13q32. Paired (on sister chromatids) signals were only seen at band 13q32. In several metaphases, signals on a single chromatid were observed at chromosome bands 6p21 or 4q21, suggesting hybridization to distantly related sequences.

EXAMPLE II

Isolation of MOAT-C and MOAT-D cDNA.

Isolation of the MOAT-B₄ transporter as described above suggested the possibility that there were other MRP/cMOAT-related transporters. A blast search (36) of the nonredundant expressed sequence tag data base using MRP and related yeast transporters revealed two clones with significant similarity to MRP and cMOAT. The first of these sequences (I.M.A.G.E. consortium clone 113196) was 1.2 kb in length, 800 bp of which encoded an MRP-related peptide. A segment of this clone was used as a probe to screen ovarian and hematopoietic bacteriophage libraries. Analysis of these cDNA clones indicated that they contained approximately 2 kb of additional coding sequence not present in clone 113196. An additional 1655 bp of 5' sequence was obtained by several rounds of RACE using the bacteriophage DNA prepared from the ovarian cDNA library as template. The continuity of the sequences obtained by RACE with the cDNA clones isolated from bacteriophage libraries was confirmed by nucleotide sequence analysis of a 2 kb product obtained by RT/PCR using an upstream oligonucleotide primer located at the 5' end of the RACE sequence and a downstream primer located at the 5' end of the cDNA obtained by plaque

hybridization. A total of approximately 5.9 kb of cDNA sequences were isolated. Nucleotide sequence analysis revealed an open reading frame of 4311 bp that was preceded by an in frame stop codon located at positions -93 (relative to the putative initiation codon), and encoding a predicted protein of 1437 amino acids, which is designated MOAT-C herein. The open reading frame was followed by approximately 1.4 kb of 3' untranslated sequences in which a polyadenylation sequence (AAUAAA) was located 20 bp upstream of the poly(A) tail. The most upstream ATG in the open reading frame was located in the sequence context "GAAGATGA". The A at position -3 of the putative translation initiation codon was in agreement with the major feature of the Kozak consensus sequence, but the A at position +4 was divergent from the more usual G (37). The second sequence identified in our data base search (I.M.A.G.E. consortium clone 208097) was 1.2 kb in length, of which 588 bp encoded an MRP-related peptide. A segment of this clone was used as a probe to screen liver and monocyte bacteriophage cDNA libraries, and 5' cDNA segments of the isolated cDNA clones were used in a subsequent round of screening. Together approximately 5.2 kb of cDNA sequence were isolated. Nucleotide sequence analysis revealed an open reading frame of 4570 bp, which is designated MOAT-D herein. The open reading frame was followed by approximately 0.6 kb of 3' untranslated sequences in which a polyadenylation sequence (AAUAAA) was located 12 bp upstream of the poly(A) tail. An upstream in frame stop codon was not present in the MOAT-D cDNA clones, and attempts to obtain additional upstream sequences by RACE using as template cDNA prepared from sources in which MOAT-D is abundant were not successful. The most upstream ATG in the open reading frame

(nucleotide position 5-7), located in the sequence context "ATGGATGG", was therefore designated as the translational initiation site. The G at position +4, was in good agreement with the Kozak consensus sequence, but the T at -3 was divergent from the more usual A (37). Although an upstream in frame stop codon was not identified in the MOAT-D cDNA clones, the size of the encoded protein was within one amino acid of the size of the transporter with which it shares the highest degree of identity (MRP), suggesting that the complete MOAT-D open reading frame was present in the isolated cDNA clones.

Analysis of the MOAT-C and MOAT-D Predicted Proteins.

Comparison of the MOAT-C and MOAT-D predicted proteins with complete coding sequences in protein data bases using the BLAST program indicated that they shared significant similarity with several eukaryotic ABC transporters. Typical features of eukaryotic ABC transporters were present in the predicted proteins. See Figure 5. Overall the proteins were composed of hydrophobic domains containing potential transmembrane spanning helices and two nucleotide binding folds. Conserved Walker A and B ATP binding sites, as well as a conserved C motif, the signature sequence of ABC transporters, was present in the nucleotide binding folds. Computer assisted analysis of potential transmembrane helices of MOAT-C using the TMAP program (19) predicted 12 transmembrane helices with 6 transmembrane spanning helices in each of two membrane spanning domains. This 6 + 6 (TM1-TM6 and TM7-TM12) configuration of predicted transmembrane helices is in agreement with topological models proposed for several other ABC transporters (20, 21), and is shown in Figure 5. However, alternative

predictions of transmembrane segments were obtained using different program parameters or input sequence alignments. Comparison of the hydropathy profiles of MOAT-C with other MRP/cMOAT-related transporters (Fig. 6B) indicates that its structure is similar to that of MOAT-B, which also has two membrane spanning domains.

In contrast to MOAT-C, hydrophobicity analysis of MOAT-D indicated that it has three membrane spanning domains. Similar to MRP, cMOAT and the yeast cadmium resistance factor 1 (YCF1), MOAT-D has an additional N-terminal hydrophobic domain that is not present in MOAT-B or MOAT-C (Figs. 5 and 6). A 5+6+6 configuration of transmembrane spanning helices has been proposed for MRP (38), in which the N-terminal extension harbors 5 transmembrane spanning helices, and 6 transmembrane helices are present in the second and third membrane spanning domain. An alignment of the MOAT-D predicted protein with MRP using the GAP program indicated that proposed MRP transmembrane spanning helices were conserved in MOAT-D. This 5+6+6 model for MOAT-D is shown in Fig. 5. Another configuration of transmembrane spanning helices (5+6+4) was predicted using computer assisted analysis. MRP has been reported to have two N-linked glycosylation sites in its N-terminus (Asn-19 and Asn-23) and another site located between the first and second transmembrane spanning helix of its third membrane spanning domain (Asn-1006). The alignment of MOAT-D with MRP indicated that an N-terminal (Asn-21) and a distal N-glycosylation sites (Asn-1008/1009) were conserved in analogous positions in MOAT-D. Only the distal N-glycosylation site of MRP is conserved in MOAT-C (Asn890) (Fig. 5) and MOAT-B⁺ (Asn746/754).

Among ABC transporters, the degree of similarity of

the nucleotide binding folds is considered to be the best indicator of functional conservation. Comparison of the nucleotide binding folds of MOAT-C and MOAT-D with other eukaryotic ABC transporters indicated that they were most closely related to those of human MRP, human cMOAT and yeast YCF1, three transporters that have organic anions as substrates. As shown in Table 2, among the human transporters, the MOAT-C NBF1 was about equally related to MOAT-D, MRP and cMOAT (55-61% identity), and less similar to MOAT-B (49% identity).

Table II. Amino acid identity: nucleotide binding folds 1 and 2 of MRP/cMOAT sub-family members.

	MOAT-C	MOAT-D	MOAT-B	MRP	cMOAT	YCF1
	%IDENTIFY (BNF1/NBF20)					
MOAT-C	-----	57.3/58.9	49.3/59.1	60.0/59.4	61.3/60.6	55.3/58/8
MOAT-D	57.3/58/9	-----	55.3/54.1	70.1/73.8	67.3/70.0	52.7/61.3
MOAT-B	49.3/59.1	55.3/54.1	-----	57.3/61/6	53.3/55.3	56.0/57.2
MRP	60.0/59.4	70.7/73.7	57.3/61.6	-----	66.0/73.1	53.3/63.8
cMOAT	61.3/60.6	67.3/70.0	53.3/55.3	66.0/73.1	-----	50.7/61/3
YCF1	55.3/58.8	52.7/61.3	56.0/57.2	53.3/63.8	50.7/61.3	-----

The MOAT-C NBF2 shared about equal amino acid identity with the five other transporters in this group (59-61% identity). Overall, the MOAT-C protein was about equally related to the other five transporters in this group, with 33.1-36.5% identity. Aside from these

transporters, MOAT-C is most closely related to CFTR, with which its NBFs shared 44%/42 % identity, and SUR, with which its NBFs shared 49%/51% identity.

The MOAT-D NBFs were clearly most closely related to those of MRP and cMOAT, with which they shared considerable amino acid identity (67.3-73.8%). See Table III. Of the latter two transporters, the MOAT-D NBFs were slightly more related to those of MRP. In contrast, the MOAT-D NBFs shared only 55.3-58.9% identity with those of MOAT-C and MOAT-B. Overall, MOAT-D was again most closely related to MRP (57.3%) and cMOAT (46.9%), but significantly more related to MRP. Consistent with the analysis of NBFs, MOAT-D was much less related to MOAT-C and MOAT-B, with which it shared only 33.1% and 35.3% identity, respectively. Alignment of the MOAT-C and MOAT-D nucleotide binding folds with those of other eukaryotic transporters is shown in Fig. 6.

Table III. Overall amino acid identifying among MRP/cMOAT sub-family members

	MOAT-C	MOAT-D	MOAT-B	MRP	cMOAT	YCF1
	%identity					
MOAT-C	----	33.1	36.5	35.8	36.2	33.6
MOAT-D	33.1	----	35.3	57.3	46.9	38.1
MOAT-B	36.4	35.3	----	39.4	36.8	38.8
MRP	35.8	57.3	39.4	----	48.4	46.4
cMOAT	36.3	46.9	36.8	48.8	----	38.8
YCF1	33.6	38.1	38.8	40.4	38.8	----

Expression Pattern of MOAT-C and MOAT-D in Human Tissues.

To gain insight into the possible functions of MOAT-C and MOAT-D, their expression patterns in a variety of human tissues was examined by RNA blot analysis. As

shown in Fig. 7 (upper panels), a MOAT-C transcript of approximately 6.6 kB was readily detected in several tissues. MOAT-C transcript levels were highest in skeletal muscle, with intermediate levels in kidney, testes, heart and brain. Low levels were detected in most other tissues, including spleen, thymus, prostate, ovary, and placenta. Prolonged exposures were required for detection in lung and liver. MOAT-D was expressed as an approximately 6 kb transcript (middle panels). Compared to MOAT-C, the MOAT-D expression pattern was more restricted. MOAT-D was highly expressed in colon and pancreas, with lower levels in liver and kidney. Low levels were detected in small intestine, placenta and prostate. Prolonged exposures were required to detect MOAT-D in testes, thymus, spleen and lung.

Chromosomal localization of the MOAT-C and MOAT-D genes.

The MOAT-C and MOAT-D chromosomal localizations were determined by fluorescence *in situ* hybridization. As shown in Figure 8, hybridization of the MOAT-C probe to metaphase spreads revealed specific labeling at human chromosome band 3q27. Fluorescent signals were detected on chromosome 3q in each of 22 metaphase spreads scored. Of 75 signals observed, 43 (57%) were on 3q. Paired (on sister chromatids) signals were only seen at band 3q27. Hybridization of the MOAT-D probe revealed specific labeling at human chromosome band 17q21.3. Fluorescent signals were detected on chromosome 17 in each of 21 metaphase spreads scored. Of 83 signals observed, 34 (41%) were on 17q21.3. Paired (on sister chromatids) signals were only seen at band 17q21.3.

EXAMPLE III**Isolation of MOAT-E and MOAT-E cDNA.**

Analysis of ara, a reported cDNA sequence that encodes a 453 amino acid transporter, revealed that it is a non-physiological sequence representing a combination of 5' MRP sequences fused to an MRP/cMOAT-related transporter. The MRP sequences extend to codon 8 of the reported predicted protein.

To isolate the complete physiological cDNA, a RT/PCR approach was employed in which primers were designed based upon a reported genomic sequence that encodes exons identical to the reported ara sequence. The MOAT-E cDNA was isolated in three segments. The first segment, spanning residues 1-616, was isolated by PCR using 5' primer ATGGCCGCGCCTGCTGAGC; (SEQ ID NO: 10) and 3' primer GTCTACGACACCAGGGTCAA (SEQ ID NO: 11). The second segment, spanning residues 1815-3187, was isolated by PCR using 5' CTGCCTGGAAGAAGTTGACC (SEQ ID NO: 12) and 3' primer CTGGAATGTCCACGTCAACC (SEQ ID NO: 13). The third segment, spanning residues 3158-1503, was isolated by PCR using 5' primer GGAGACAGACACGGTTGACG (SEQ ID NO: 14) and 3' primer GCAGACCAGGCCTGACTCC (SEQ ID NO: 15). The primer were designed based upon the nucleotide sequence of human genomic BAC clone CIT987SD-962B4. The template for these reactions was random-primed human kidney cDNA prepared from total RNA. Using this approach the physiological cDNA was isolated which is designated MOAT-E herein and set forth as Sequence I.D. No. 7.

Analysis of the MOAT-E Predicted Protein.

MOAT-E encodes a 1503 amino acid transporter. The MOAT-E predicted amino acid sequence is designated Sequence I.D. No. 8. See Figure 9. Also shown is the

location of potential transmembrane helices (overbars), potential N-glycosylation site (black dot) and the two nucleotide binding folds (NBF1 and NBF2). Walker A and B motifs, as well as the signature C motif of ABC transporters are also indicated. Comparison of MOAT-E with ara indicates that the ara predicted protein is not only a fused sequence, but also that it represents only 446 (~30%) of the 1503 MOAT-E residues.

Comparison of MOAT-E with the other members of the MRP/cMOAT subfamily, which include MRP, cMOAT, MOAT-B, MOAT-C and MOAT-E, is shown in Table IV. MOAT-E is highly related to MOAT-D, MRP and cMOAT, with which it shares 39-45% identity. This high degree of identity is also indicated by the high percent identities of the nucleotide binding folds, which range from 55-61%. In contrast, MOAT-E is less related to MOAT-B and MOAT-C, with which it shares ~31% and 34% identity, respectively.

Table IV. Amino acid identity among MRP/cMOAT sub-family members.^a The bold type indicates the percent identity of the overall proteins, and the parentheses indicates the percent identity of the nucleotide binding folds.

	MOAT-E	MOAT-B	MOAT-C	MOAT-D	MRP	cMOAT
	% identity ^b					
MOAT-E	---	33.9	30.6	43.6	45.1	38.9
	---	(52.0/56.6)	(50.0/52.5)	(59.3/59.4)	(61.3/61.4)	(55.3/59.4)
MOAT-B	33.9	---	36.4	35.3	39.4	36.8
	(52.0/56.6)	---	(49.3/59.1)	(55.3/54.1)	(57.3/61.6)	(56.0/57.2)
MOAT-C	30.0	36.4	---	33.1	35.8	36.2
	(50.0/52.5)	(49.3/59.1)	---	(57.3/58.9)	(60.6/59.4)	(61.3/60.6)
MOAT-D	43.6	35.3	33.1	---	57.3	46.9
	(59.3/59.4)	(55.3/54.1)	(57.3/58.9)	---	(70.7/73.8)	(67.3/70.0)
MRP	45.1	39.4	35.8	57.3	---	48.4
	(61.3/61.9)	(57.3/61.6)	(60.0/59.4)	(70.7/73.8)	---	(66.0/73.1)
cMOAT	38.9	36.8	36.2	46.9	48.4	---
	(53.1/59.4)	(56.0/57.2)	(61.3/60.6)	(67.3/70.0)	(66.0/73.1)	---

^aoverall amino acid identities are indicated in bold-face, and identities of nucleotide binding folds 1 and 2 are indicated in parentheses (NBF1/NBF2).
^bpercent identity was obtained using the GAP command in the GCG package.

Comparison of the hydropathy profile of MOAT-E with other members of the MRP/cMOAT subfamily is shown in figure 10. The data reveal that MOAT-E has a hydrophobic N-terminal segment that is present in its closest relatives, MOAT-D, MRP and cMOAT. This structural feature is present in all of the currently known organic anion transporters, and suggests that MOAT-E may share substrate specificity with MRP and cMOAT. MOAT-E may also share the drug resistance activity of the latter two proteins. In contrast, MOAT-B and MOAT-C do not have this hydrophobic N-terminal extension.

Expression Pattern of MOAT-E in Human Tissues.

In a Northern blot of RNA isolated from various tissues, MOAT-E expression is restricted to liver and kidney, suggesting that MOAT-E may participate the excretion of substances into the urine and bile. See Figure 11. This figure also shows that MOAT-E is expressed as an ~6 kB transcript. This is in contrast to the ~2.3 kB transcript that was reported for ara, clearly indicating that the fused ara transcript is unique to the cell line from which it was isolated, and is not a physiological transcript. Together, the isolation of MOAT-E and analysis of its sequence and expression pattern suggest that it may be involved in cellular resistance to drugs and/or the excretion of drugs into the urine and bile.

DISCUSSION

The present invention discloses additional MRP/cMOAT-related transporters which were identified by

using a degenerative PCR cloning approach in which the conserved amino terminal ATP-binding domain of known eukaryotic transporters was targeted. Using this approach the complete coding sequences of MOAT-B, MOAT-C, MOAT-D and MOAT-E were obtained. MOAT-B is a protein whose predicted structure indicates that it is a member of the ABC transporter family. Comparison of the MOAT-B predicted protein with other transporters reveals that it is most closely related to MRP, cMOAT and yeast YCF1, and thus extends the number of known full length MRP-related transporters. The similarity of MOAT-B to these transporters suggest that it shares a similar substrate specificity. Transport assays using membrane vesicle preparations indicate that MRP is capable of transporting diverse organic anions, including glutathione S-conjugates such as LTC₄, oxidized glutathione, and glucuronidated and sulfated conjugates of steroid hormones and bile salts (7). Although membrane vesicle transport assays of substrate specificity using cMOAT-transfected cells have not yet been reported, genetic and biochemical studies using TR- and EHBR rat strains, which are defective in the hepatobiliary excretion of glutathione and glucuronate conjugates, indicate that it is also an ATP-dependent transporter of organic anions. cMOAT, which is primarily expressed in the canalicular membrane of hepatocytes, has been reported to be absent in these rat strains, and hepatocyte canalicular membranes prepared from the mutant rats are deficient in the ATP-dependent transport of glutathione and glucuronate conjugates (23, 24). In addition, cMOAT protein has also been reported to be absent in the hepatocytes of patients with Dubin-Johnson syndrome (25), a disorder manifested by chronic

conjugated hyperbilirubinemia. YCF1, a yeast transporter, has also been demonstrated to transport glutathione complexes (26). Thus, based upon the similarity of MOAT-B to these three transporters, it is possible that it also functions to transport organic anions, an activity critical to the cellular detoxification of a wide range of xenobiotics.

MOAT-C, MOAT-D and MOAT-E are three other MRP/cMOAT-related transporters. The isolation of these two transporters extends the number of known full length members of this subfamily to six. Based upon the degree of amino acid similarity and overall topology these six proteins fall into two groups. The first group is composed of MOAT-D, MOAT-E, MRP and cMOAT. These four transporters are highly related, sharing ~39-45% amino acid identity. MOAT-D is more closely related to MRP (57% identity) than is cMOAT (48% identity), and is therefore the closest known relative of MRP. In addition to a high degree of amino acid identity, the similarity between MOAT-D, MRP and cMOAT, also extends to overall topology. Like MRP and cMOAT, MOAT-D and MOAT-E have three membrane spanning domains, including an N-terminal hydrophobic extension that is predicted to harbor ~5 transmembrane helices, and which is absent in transporters such as CFTR and MDR1. This N-terminal extension is also present in YCF1, a related yeast transporter that transports glutathione S-conjugates, and SUR, a more distantly related transporter involved in the regulation of potassium channels. The second group of MRP/cMOAT-related transporters is composed of MOAT-B and MOAT-C. These two transporters are distinguished from the first group by their lower level of amino acid similarity and distinct topology. Like MOAT-D and MOAT-E, MOAT-B

and MOAT-C are more closely related to MRP (39% and 36%, respectively) and cMOAT (37% and 36%, respectively) than to other eukaryotic transporters. However, they share considerably less similarity with MRP, cMOAT, MOAT-D and MOAT-E than the latter four transporters share with each other (~39-45% identity). In addition, in contrast to MRP, cMOAT, MOAT-D and MOAT-E, MOAT-B and MOAT-C do not have an N-terminal membrane spanning domain, and their topology is therefore more similar to many other eukaryotic ABC transporters that also have only two membrane spanning domains.

Defining the contributions of MOAT-B, MOAT-C, MOAT-D and MOAT-E to cytotoxic drug resistance will facilitate the design of novel chemotherapeutic agents. The multidrug resistance activity of MRP is well described. While the drug sensitivity pattern of cMOAT-transfected cells has not yet been reported, the possibility that it may also confer resistance to cytotoxic drugs is suggested by a recent report in which transfection of a cMOAT antisense vector was found to enhance the sensitivity of a human liver cancer cell line to both natural product drugs and cisplatin. Since MOAT-D and MOAT-E are more closely related to MRP than is cMOAT, the possibility that they will also confer resistance is particularly intriguing. The availability of the MOAT-B, MOAT-C, MOAT-D and MOAT-E cDNAs will facilitate the analysis of their possible contributions to cytotoxic resistance.

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While certain of the preferred embodiments of the present invention have been described and specifically exemplified above, it is not intended that the invention be limited to such embodiments. Various modifications may be made thereto without departing from the scope and spirit of the present invention, as set forth in the following claims.

What is claimed is:

1. An isolated nucleic acid molecule having the sequence of SEQ ID NO:1, said nucleic acid molecule comprising a nucleotide sequence encoding a MOAT-B transporter protein about 1350 amino acids in length, said encoded transporter protein comprising a multi-domain structure including a tandem repeat of nucleotide binding folds appended C-terminal to a hydrophobic domain, said nucleotide binding folds having Walker A and B ATP binding sites, said C-terminal domain having a plurality of membrane spanning helices.
2. The nucleic acid molecule of claim 1, which is DNA.
3. The DNA molecule of claim 2, which is a cDNA comprising a sequence approximately 5.9 kilobase pairs in length that encodes said MOAT-B transporter protein.
4. The DNA molecule of claim 2, which is a gene comprising introns and exons, the exons of said gene specifically hybridizing with the nucleic acid of SEQ ID NO 1, and said exons encoding said MOAT-B transporter protein.
5. An isolated RNA molecule transcribed from the nucleic acid of claim 1.
6. The nucleic acid molecule of claim 1, wherein said sequence encodes a MOAT-B transporter

protein having an amino acid sequence selected from the group consisting of SEQ ID NO 2 and amino acid sequences encoded by natural allelic variants of said sequence.

7. The nucleic acid molecule of claim 6, which comprises SEQ ID NO 1.

8. An antibody immunologically specific for the protein encoded by the nucleic acid of claim 1.

9. An antibody as claimed in claim 8, said antibody being monoclonal.

10. An antibody as claimed in claim 8, said antibody being polyclonal.

11. An isolated nucleic acid molecule having the sequence of SEQ ID NO: 3, said nucleic acid molecule comprising a sequence encoding a MOAT-C transporter protein about 1450 amino acids in length, said transporter protein having a multi-domain structure including a tandem repeat of nucleotide binding folds, said nucleotide binding folds having Walker A and B binding sites, and a C-terminal hydrophobic domain that contains several membrane spanning helices.

12. The nucleic acid molecule of claim 11, which is DNA.

13. The DNA molecule of claim 12, which is a cDNA comprising a sequence approximately 6.6 kilobase pairs in length that encodes said MOAT-C transporter protein.

14. The DNA molecule of claim 12, which is a gene comprising introns and exons, the exons of said gene specifically hybridizing with the nucleic acid of SEQ ID NO 3, and said exons encoding said MOAT-C transporter protein.

15. An isolated RNA molecule transcribed from the nucleic acid of claim 11.

16. The nucleic acid molecule of claim 11, wherein said sequence encodes a MOAT-C transporter protein having an amino acid sequence selected from the group consisting of SEQ ID NO 4 and amino acid sequences encoded by natural allelic variants of said sequence.

17. The nucleic acid molecule of claim 11, which comprises SEQ ID NO 3.

18. An antibody immunologically specific for the protein encoded by the nucleic acid of claim 11.

19. An antibody as claimed in claim 18, said antibody being monoclonal.

20. An antibody as claimed in claim 18, said antibody being polyclonal.

21. An oligonucleotide between about 10 and about 200 nucleotides in length, which specifically hybridizes with a protein translation initiation site in a nucleotide sequence encoding amino acids of SEQ ID NO 4.

22. An oligonucleotide between about 10 and about 200 nucleotides in length, which specifically hybridizes with a protein translation initiation site in a nucleotide sequence encoding amino acids of SEQ ID NO 2.

23. An isolated nucleic acid molecule having the sequence of SEQ ID NO: 5, said nucleic acid molecule comprising a sequence encoding a MOAT-D transporter protein about 1550 amino acids in length, said transporter protein having a multi-domain structure including a tandem repeat of nucleotide binding folds, said nucleotide binding folds having Walker A and B binding sites, and a C-terminal hydrophobic domain that contains several membrane spanning helices.

24. The nucleic acid molecule of claim 23, which is DNA.

25. The DNA molecule of claim 24, which is a cDNA comprising a sequence approximately 6 kilobase pairs in length that encodes said MOAT-D transporter protein.

26. The DNA molecule of claim 24, which is a gene comprising introns and exons, the exons of said gene specifically hybridizing with the nucleic acid of SEQ ID NO 5, and said exons encoding said MOAT-D transporter protein.

27. An isolated RNA molecule transcribed from the nucleic acid of claim 23.

28. The nucleic acid molecule of claim 23, wherein

said sequence encodes a MOAT-D transporter protein having an amino acid sequence selected from the group consisting of SEQ ID NO 6 and amino acid sequences encoded by natural allelic variants of said sequence.

29. The nucleic acid molecule of claim 23, which comprises SEQ ID NO 5.

30. An antibody immunologically specific for the protein encoded by the nucleic acid of claim 23.

31. An antibody as claimed in claim 30, said antibody being monoclonal.

32. An antibody as claimed in claim 30, said antibody being polyclonal.

33. An oligonucleotide between about 10 and about 200 nucleotides in length, which specifically hybridizes with a protein translation initiation site in a nucleotide sequence encoding amino acids of SEQ ID NO 6.

34. An isolated nucleic acid molecule having the sequence of SEQ ID NO:7, said nucleic acid molecule comprising a nucleotide sequence encoding a MOAT-E transporter protein about 1503 amino acids in length, said transporter protein having a multi-domain structure including a tandem repeat of nucleotide binding folds, said nucleotide binding folds having Walker A and B binding sites, and a C-terminal hydrophobic domain that contains several membrane spanning helices.

35. The nucleic acid molecule of claim 34,

which is DNA.

36. The DNA molecule of claim 35, which is a cDNA comprising a sequence approximately 6 kilobase pairs in length that encodes said MOAT-E transporter protein.

37. The DNA molecule of claim 35, which is a gene comprising introns and exons, the exons of said gene specifically hybridizing with the nucleic acid of SEQ ID NO 7, and said exons encoding said MOAT-E transporter protein.

38. An isolated RNA molecule transcribed from the nucleic acid of claim 34.

39. The nucleic acid molecule of claim 34, wherein said sequence encodes a MOAT-E transporter protein having an amino acid sequence selected from the group consisting of SEQ ID NO 8 and amino acid sequences encoded by natural allelic variants of said sequence.

40. The nucleic acid molecule of claim 39, which comprises SEQ ID NO 7.

41. An antibody immunologically specific for the protein encoded by the nucleic acid of claim 34.

42. An antibody as claimed in claim 41, said antibody being monoclonal.

43. An antibody as claimed in claim 41, said antibody being polyclonal.

44. An oligonucleotide between about 10 and about 200 nucleotides in length, which specifically hybridizes with a protein translation initiation site in a nucleotide sequence encoding amino acids of SEQ ID NO 7.

45. A plasmid comprising a nucleotide sequence selected from the group consisting of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5 and SEQ ID NO:7.

46. A vector comprising a nucleotide sequence selected from the group consisting of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5 and SEQ ID NO:7.

47. A retroviral vector comprising a nucleotide sequence selected from the group consisting of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5 and SEQ ID NO:7.

48. A host cell comprising at least one nucleic acid molecule having a sequence selected from the group consisting of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5 and SEQ ID NO:7.

49. A host cell as claimed in claim 48, wherein said host cell is selected from the group consisting of bacterial, fungal, mammalian, insect and plant cells.

50. A host cell as claimed in claim 48, wherein said nucleic acid is provided in a plasmid and is operably linked to mammalian regulatory elements which confer high expression and stability of mRNA transcribed from said nucleic acid.

51. A host cell as claimed in claim 48, wherein said nucleic acid is provided in a plasmid and is operably linked to mammalian regulatory control elements in reverse anti-sense orientation.

52. A host animal comprising at least one nucleic acid molecule selected from the group consisting of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5 and SEQ ID NO: 7.

53. A host animal as claimed in claim 52, wherein said animal harbors a homozygous null mutation in its endogenous MOAT gene wherein said mutation has been introduced into said mouse or an ancestor of said mouse via homologous recombination in embryonic stem cells, and further wherein said mouse does not express a functional mouse MOAT protein.

54. The transgenic mouse of claim 53, wherein said mouse is fertile and transmits said null mutation to its offspring.

55. The transgenic mouse of claim 53, wherein said null mutation has been introduced into an ancestor of said mouse at an embryonic stage following microinjection of embryonic stem cells into a mouse blastocyst.

56. A method for screening a test compound for inhibition of MOAT mediated transport, comprising:

a) providing a host cell expressing at least one MOAT-encoding nucleic acid having a sequence selected from the group consisting of SEQ ID NOS: 1, 3, 5, and 7;

b) contacting said host cell with a compound suspected of inhibiting MOAT-mediated transporter activity; and

c) assessing inhibition of transport mediated by said compound.

57. A method as claimed in claim 56, wherein inhibition of MOAT mediated transport is indicated by restoration of anticancer drug sensitivity.

58. A method as claimed in claim 57, wherein said inhibition of MOAT mediated transport is indicated by a reduction of transporter mediated cellular efflux of anticancer agents.

59. A kit for detecting the presence of MOAT encoding nucleic acids in a sample, comprising:

- a) oligonucleotide primers specific for amplification of MOAT encoding nucleic acids;
- b) polymerase enzyme;
- c) amplification buffer; and
- d) MOAT specific DNA for use as a positive control.

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MOAT-B
MRP 1 MALRGFCSADGSDPLWDMVNTWNTSNPDFTKCFONTVLVWVPCFYLWACFFPYFLYLSRHRDGYIQMTPLNKTALGFLLLWIVCWADLFYSFWERSRGI 100
MOAT-B 1MLP 3
MRP 101 FLAPVFLVSPTLLGITTLLATFLIQLERRKGVSSGIMTLFWLVALVCAILRSKIMTALKEDAQVDLFRDITFYVYFSLLLIQLVLSCFSDRSPLFSE 200
MOAT-B 4 VYQEVKPNPLQDANICSRVFFWMLNPLFKIGHKRRLEEDMYSVLPEDRSOHLGEELQGFWDKEVLRAENDAOK 77
MRP 201 TINDPNPCPESSASFLSRITFWITGLIVRGYRQPLEGSDLSLNKEDTSEQVVPVLVKNWKECAKTRKQPVKVYSSKDPAPKESKVDANEVEAL 300
MOAT-B 78PSLTRAIIKCYWKSYLVLGIFTLIEESAKVIQIFLGKIINYFENYDPMDSVALNTAYAYATVLTFTLILAILHHLYFYHVOCAGHRL 166
MRP 301 IVKSPOKEWNPFLKVLKTFPGPYFLMSFFFKATHDLHMFSGPQILKLLIKFVNDTKAPDMQGY.....FYTULLFVTACLQTLVLHLYFHICFVSGHRI 395
MOAT-B 167 RVAMCHMIYRKALRLSNMAMGKTTTGOIVNLLSNDVNKFDQVTVFLHFLWAGPLOAIJAVTALLHMEIGISCLAGHAVLIILLPQSCFGKLFSSLSRSKTA 266
MRP 396 KTAVIGAVYRKALVITNSARKSSTVGEIVNLSVDAQRFDMLATYINMWSAPLOVILALYLLWNLGSPVLGAVAVHVMVPMVAVMAMKTKTYQVAHM 495
MOAT-B 267 TFTDARIRTHNEVITGIRIIKHYAWEKSFNLTINLRKKEISKILRSSCLRGHNLASFFSASKIIVFVFTTYVLLG...SVITASRVFVAVTLYGAVRLT 364
MRP 496 KSKDNRIKLWNEILNGIKVLKYAWELAFKDKVLAIROEELKVLKKSAYLSAVGTFTWCTPFLVALCTFAVYVTIDENILDAQTAFVSLALFNILRFP 595
MOAT-B 365 VTLFFPSAIERVSEAIVSIRRIQTFLLEIS...ORNRQLPSDGKQHVVDFTAFWDKASETPTLQGLSFTVRPGELLAVVGPVGAGKSSLLSAVLG 460
MRP 596 LNI.LPMVISSIVQASVSLKRLRIFLSHEELEPDSEIRRPVKDGGTNSITVRNATFTWAR.SDPPTLNGITFSIPEGALVAVVGVQVCGKLSLLSALLA 693
MOAT-B 461 ELAPSHGLVSVHGRIAVVSQOPWVFSGLTSLRNILFGKKYKERYEKVIKACALKKDLQLEDGDLTVIGDRGTTLSGGQKARVNARAVYQDADIYLLDD 560
MRP 694 EMDKVEGHVAIKGSVAVVPPQAWIQNDSLRNIFGCGLEPPYRSVIGACALLPDLIELPSGDRTEIGKGVNLGGQKQVSLARAVSNADIYLLDD 793
MOAT-B 561 PLSAVDAEVSRLHFLCICQ...ILHEKITILVTHQLQYLKAAQOILKDKGMVOKGTYTEFLKSGIDFGSLK.....KDNEEQPPVPG..... 645
MRP 794 PLSAVDAHVGKHIFENVIGPKMLKNKTRILVTHMSYLPQVDVIVMSGGKISEHGSYQELLARDGAFAEFLRTYASTEQQEQAENGVTGVSFGPGKEA 893
MOAT-B 646TPTLRNRTFSESSVMSQSSRPSLKDGALESQDT...ENVPVTLSEENRSEKGVGFQAYKNYFRAGAHMIVFIFILLNTAAQVAVYVLO 731
MRP 894 KQMENGHLVTDASAGQLQRLSSSSSYSGDISRHNNSTAELOKAEKKEETWKLHEADKAQTQGVKLSVYWDYHKAIGLFSISFLF.MCNHVSALAS 992
MOAT-B 732 DWLWSYWANKQSHLNTVNGGGNVTEKLDLWYLGIIYSGLTATVTLFGIARSLVYVVLVNSSQTLHNMHIFESILKAPVLPFDRNPIGRILNRFKIDIGH 831
MRP 993 NYWLSLWTD...DPIVNGTQENTKVR...LSVYGALGISQGIIVFGYSMAVSGGILASRCLHVDLLHSILRSPHSFFERTPSGNLVNRFKIDELT 1082
MOAT-B 832 LDDLLPLTFLDFIQTLLQVGVVSVAVAVIPWIAIPLVPLGIIIFILRRYFLETSRDVKLESTTRSVPVSHLSSSLQGLWITRAYKAEERQELFDAHQ 931
MRP 1083 VDSMIPEVIKMFNGSLFNVIGACIVILLATPIAAIIIPPLGLIYFFVQRFYVASSRQLKRLLESVRSRSPVYSHFNETLLGVSVIRAFEEQERFIHQSDLKV 1182
MOAT-B 932 DLHSEAWFLFTTSRWFAVRLDAICAMFVIIAFAFGSLILAKTLDAQOVGLALSALTLMHGFOWCVROSAEVENMMHISVERVIEYTDLEKEAPWEYQK.R 1030
MRP 1183 DENQKATYPSIVANRWLAARLECVGNCIVLFAALFAVISRHSLSAGLVGLSVSYSLQVTTYLNLVMSSEMETNIVAVERLKEYSETEKEAPWQIQETR 1282
MOAT-B 1031 PPPAWPHEGVIIIFDNVNFMYSPGGPLVLKHLTALIKSQEKVGVIGRTGACKSSLSLALFRLSE.PEGKIWDKILTTEIGLHDLRKHMSIIPQEPVLFTG 1129
MRP 1283 PPSSWPQVGRVEFRNYCLRYREDLDFVLRHINVTINGGEKVGIVGRTGACKSSLTGLFRINESAEGEIIIDGINIAKIGLHDLRKFITIIIPQDPVLFSG 1382
MOAT-B 1130 THRNLDPPFKENTDEELWALQEVOLKETIEDLPGKMDTELAESGNSFSVQORQLVCLARAILRKNQILIDEATANVDPRTDELIQKKIREKFAHCTVL 1229
MRP 1383 SLRNLDPPSOYSDEEVWTSLELAHLKDFVSALPKDLOHECAEGENLSVGQORQLVCLARALLRKTKILVLEATAAVDLETDLIQSTIRTOFEDCTVL 1482
MOAT-B 1230 TIAHRLNTIIDSOKIMVLDSCRLKEYDEPVYLLQNKESLFYKHYVQQLGKAEAAALTETAKQVYFKRNYHIGHTDHVNTNSNGQPSTLTIFETAL 1325
MRP 1483 TIAHRLNTIMDYTRVIVLDKGEIQEYGAPSDLLQOR.GLFYSMAKDAGLV 1531

Figure 1

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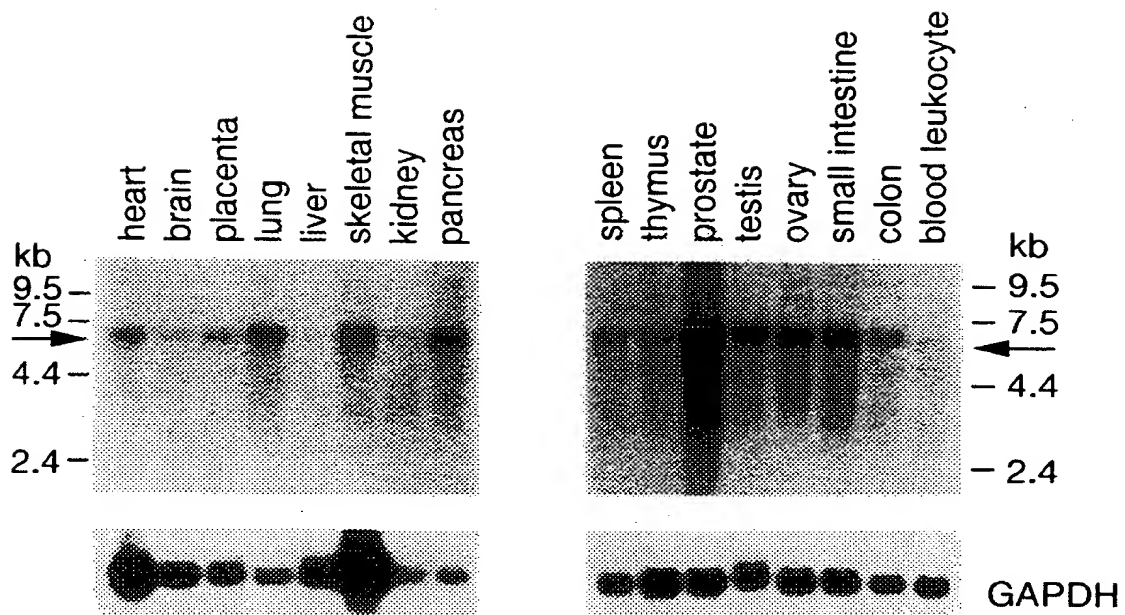


Figure 3

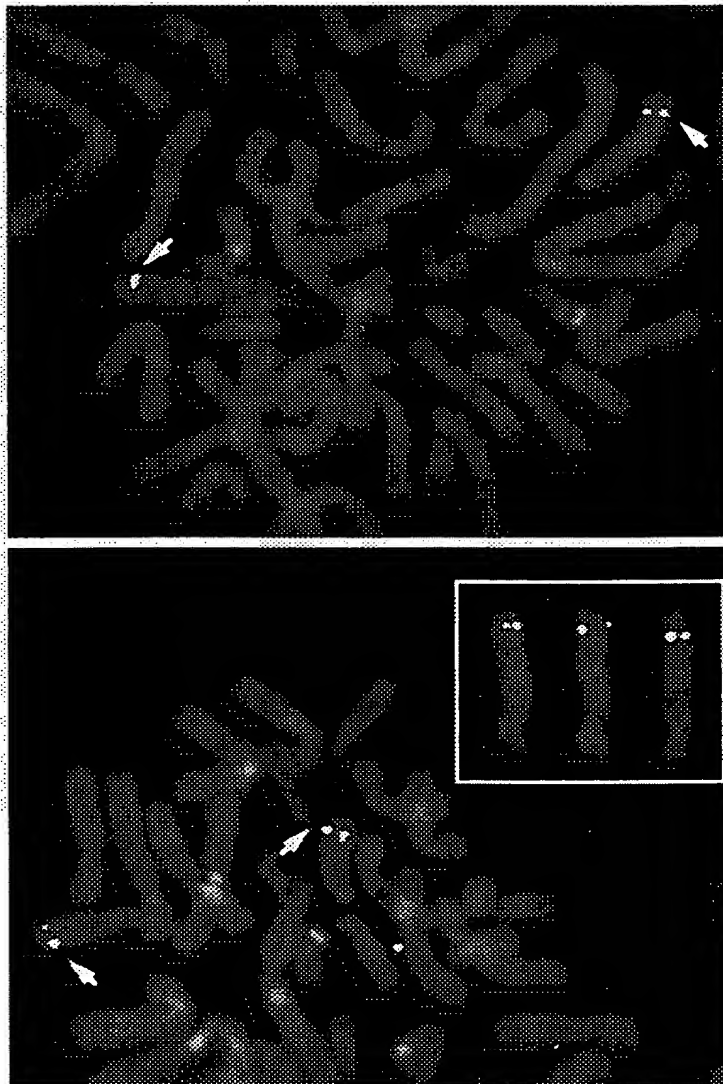


Figure 4

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Fig. 5A

1 MKDIDIGKEY IIPSPGYRSV RERTSTSGTH RDREDSKFRR TRPLECQDAL ETAARAEGLS
61 LDASMHSQLR ILDEEHPK GK YHGLSALKP IRTTSKHQHP VDNAGLFSCM TFSWLSSLAR
121 VAHKKGELSM EDVWSLSKHE SSDVNCRRLE RLWQEELNEV GPDAASLRRV VWIFCRTRLI
TM1 TM2
181 LSIVCLMITQ LAGFSGPAPM VKHLLLEYTQA TESNLQYSL LVLGLLLTEI VRSWSLALTW
TM3
241 ALNYRTGVRL RGAILTMAFK KILKLNKIE KSLGELINIC SNDGQRMFEA AAVGSLLAGG
TM4
301 PVVAILGMIY NVIILGPTGF LGSVFILFY PAMMFASRLT AYFRRCVAA TDERVQKME
TM5
361 VLTYYKFIKM YAWVKAFSQS VQKIREEERR ILEKAGYFQG ITVGVAPIVV VIASVVTFSV
TM6
421 BMTLGFDLTA AQAFVTVTVF NSMTFALKVT PFSVKSLSEA SVAVDRFKSL FLMEEVHMIK
481 NKPASPHIKI EMKNATLAWD SSHSSIQNSP KLTPKMKKDK RASRGKKEKV RQLQRTHEQA
541 VLAEQKGHL L LDSERPSPE EEEGKHIHLG HLRLQRTLHS IDLEIQEGKL VGICGSVSGS
NBFI
601 KTSLSAILG QMTLEGSIA ISGTFAYVAQ QAWILNATLR DNILFGKEYD EERYNSVLNS
A
661 CCLRPDLAIL PSSDLTEIGE RGANLSCGQR ORISLARALY SDRSIYILDD PLSALDAHVG
NBFI
721 NHIFNSAIRK HLKSKTVLFV THQLQYLVD C DEVIFMKEGC ITERGTHEEL MNLNGDYATI
B
781 FNNLLGETP PVEINSKKT SGSQKKSQDK GPKTGSVKKE KAVKPEEGQL VQLEEKQGS
TM7
841 VPWSVYGVYI QAAGGPLAF L VIMALFHLNV GSTAFSTWWL SYWIKQSGN TTVTRGNETS
TM8
901 VSDSMKDNPH MQYASIIAL SMAVMLILKA IRGVVFVKGT LRASSRLHDE LFERRILRSPM
TM9
961 KFFDTTPTGR ILNRFSKDMD EVDVRLPFA EMFIQNVILV FFCVGMHAGV FPWFLVAVGP
TM10
1021 LVILFSVLHI VSRVLIRELK RLDNITOSPF LSHITSSIQG LATIHAYNKG QEFLERYQEL
TM11 TM12
1081 LDDNQAPFFL FTCAMRWLAV RLDLISIALI TTTGLMTVLM HGQIPPAYAG LAISTAVOLT
1141 GLFQFTVRLA SETEARFTSV ERINHYIKTL SLEAPARIKN KAPSPDWPQE GEVTFENAEM
NBFI
1201 RYRENLPVLV KVSFTIKPK EKIGIVGRTC SGKSSIGMAL FRLVELSGGC IKIDGVRISD
A
1261 IGLADLRSLK SIIPQEPVLF SGTVRSNLDP FNQYTEDQIW DALERTHMKE CIAQLPLKLE
NBFI
1321 SEVMENGDNF SVGERQLLCI ARALLRHCKI LILDEATAAM DTETDL LIQE TIREAFADCT
C B
1381 MTIAHRLHT VLGSDRIMVL AQGQVVEFDT PSVLLSNDSS RFYAMFAAAE NKVAVKG

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Fig. 5B

1 MGPMDALCGS GELGSKFWD^{*} NLSVHTENPD LTPCFQNSLL AWPVCIYLWV ALPCYLLYL^{TM1}
 61 HHCRGYIILS HLSKLMVLG VLLWCVSWAD LFYSFHGLVH GRAPAPVFFV TPLVVGVTML^{TM2}
 121 LATLLIQYER LQGVQSSGVL IIFWFLCVVC AIVPFRSKIL LAKAEGEISD PFRFTTFYIH^{TM3}
 181 FALVLSALIL ACFREKPPFF SAKNVDPNPY PETS^{TM4}VGFLSR LFFWWFTKMA IYGYRHPLEE^{TM5}
 241 KDLWSLKEED RSQMVVOQLL EAWRKQEKQT ARHKASAAPG KNASGEDEV^{TM6}L LGARPRPRKP
 301 SFLKALLATF GSSFLISACF KLIQDLLSFI NPQLLSILIR FISNPMAPSW WGFLVAGLMF^{TM7}
 361 LCSMMQSLIL QHYHYIFVT GVKFRTGIMG VIYRKALVIT NSVKRASTVG EIVNLM^{TM8}SVDA^{TM9}
 421 QRFMDLAPFL NLLWSAPLOI ILAIYFLWON LGPSVLAGVA FMVLLIPLNG AVAVK^{TM10}MRAFO^{TM11}
 481 VKQMKLKDSR IKLMSEILNG IKVLKLYAWE PSFLKQVEGI ROGE^{TM11}LQLLRT AAYLHTTTT^{NBF1}
 541 TWMCS^APF^BLVT LITLWVYVYV DPNNVLDAEK AFVS^CVS^BLFNI LRLPLNMLPQ LISNLTQASV
 601 SLKRIQQFLS QEELDPQSVE RKTISPGYAI TIHSGTFTWA QDL^{NBF1}PPTLHSL DIQVPKGALV
 661 AVVGPVGC^AGK SSLVSALLGE MEKLEGKVHM KGSVAYVPOQ AWIQNCTLOE NVLFGKALNP
 721 KRYQOTLEAC ALLADLEMLP GGDQTEIGEK GINLSGGQ^CRO RVSLARAVYS DADIFLLDDP^B
 781 LSAVD^{NBF1}SHVAK HIFDEHIGPE GVLAKTRVL VTHGISFLPQ TDFIIVLADG QVSEMGYPYA
 841 LLQRNGSFAN FLCNYAPDED QGHLEDSWTA LEGAEDKEAL LIEDT^BLSNHT DLTDNDPVTY
 901 VVQKQFMRQL SALSSDGEQ GRPVPRRHG PSEKVQVTEA KADGALTQEE KAAIGTVELS
 961 VFWDYAKAVG LCITLAICLL YVGOSAAAIG ANVWLSAWTN DAMADSRQNN TSLRLGVYAA^{TM12}
 1021 LGILQGF^{TM13}LVH LAAMAAAGG IQAARVLEQA LLENKIRSPQ SFFDTTPSGR ILNCF^{TM14}SKDIY^{TM15}
 1081 VVDEV^{TM14}LAPVI LMLLNSFFNA ISTLVVIMAS TPLFTVVILP LAVLYTLVOR FYAATSRQLK
 1141 RLESVSRSP^{TM16}I YSEFSETVTG ASVIRAYNRS RDFEII^{TM17}SDTK V^{NBF2}DANQRSCYP YIISNRWLSI
 1201 GVEFVGNCVV LFAALFAVIG RSSLN^{TM17}PGLVG LSVSYSLQVT FALNWMIRMH SDLESNIVAV
 1261 ERVKEYSKTE TEAPWVVEGS RPPEGWPPRG EVEFRNYSVR YRPGDLVL^{NBF2}LR DLSLHVEGGE
 1321 KVGIVGRTGA GKSSMTLC^ALF RILEAAKGEI RIDGLNVADI GLHDLRSQLT IIPQDPILFS
 1381 GTLRMNLDPF GSYSEEDIW ALELSHLETF VSSQ^{NBF2}PAGLDF QCSEGGENLS VGORQLVCLA^C
 1441 RALLRKS^BRIL VLDEATAAID LETDNLIQAT IRTQFDTCTV LTIAHRLNTI MDYTRVLVLD
 1501 KGVVAEFDSP ANLIAARGIF YGMARDAGLA

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Nucleotide Binding Fold I									
MOAT-D	CHSLDTQVPK	CAVAVVQPV	SCGKSSVSA	LPPEYEKIE	K.....	HM	KESVAVVFN	AKSCSLCE 707
MRP	NCPTFSPE	CAVAVVQPV	SCGKSSLSA	LAENDKVEG	H.....	AI	KESVAVVFN	AKSCSLCE 724
cMOAT	VROVNDLMA	CAVAVVQPV	SCGKSSLSA	MAEENKVEG	HI.....	TI	KETTVVFN	AKSCSLCE 717
MOAT-C	VHSIDLEQE	CAVAVVQPV	SCGKSSLSA	LAENDKVEG	ST.....	AI	SETTVVFN	AKSCSLCE 717
MOAT-B	ICGLSFTVRP	CAVAVVQPV	SCGKSSLSA	VGLAPSHG	SV	HRIVVFN	AKSCSLCE 491
CFTR	IKLNFYPER	COLLAFAST	SCGKSSLSA	ITGLEPSG	KH	SRISFC	AKSCSLCE 504
SUR	ISNFTIRPR	COLLAFAST	SCGKSSLSA	ALAEKQVSE	AFWSSLPDS	EIGEDRSPER	ETATDLDIRK	RSPVAAK	AKSCSLCE 785
MDR1	IKGLNKKVQS	COLLAFAST	SCGKSSLSA	MORLYDPTG	MSVDSQDIR	TINVRFLREI	IGVVE	AKSCSLCE 486
A									
MOAT-D	NVLFCKA.LN	PRRYQOTLEA	CALLADPEML	EGGCEIGE	KSNLSGGOR	QVSLARAVY	SNADIFLDD	PLSAVDAHA	KHIDHW 793
MRP	NILFGCO.LE	PRRYQOTLEA	CALLADPEML	EGGCEIGE	KSNLSGGOR	QVSLARAVY	SNADIFLDD	PLSAVDAHA	KHIDHW 810
cMOAT	NILFGTE.FN	PRRYQOTLEA	CALLADPEML	EGGCEIGE	KSNLSGGOR	QVSLARAVY	SNADIFLDD	PLSAVDAHA	KHIDHW 803
MOAT-C	NILFGKE.YD	PRRYQOTLEA	CALLADPEML	EGGCEIGE	KSNLSGGOR	QVSLARAVY	SNADIFLDD	PLSAVDAHA	KHIDHW 727
MOAT-B	NILFGKK.YE	PRRYQOTLEA	CALLADPEML	EGGCEIGE	KSNLSGGOR	QVSLARAVY	SNADIFLDD	PLSAVDAHA	KHIDHW 577
CFTR	NILFVS.YD	PRRYQOTLEA	CALLADPEML	EGGCEIGE	KSNLSGGOR	QVSLARAVY	SNADIFLDD	PLSAVDAHA	KHIDHW 590
SUR	NILFESP.FN	PRRYQOTLEA	CALLADPEML	EGGCEIGE	KSNLSGGOR	QVSLARAVY	SNADIFLDD	PLSAVDAHA	KHIDHW 871
MDR1	IRRYRENVT	MDEIEKAVKE	ANAYDFIMK	SHKFDLVL	PSACSSGGK	PSAINGLV	RNPKLLE	ATLITTESE	AVVQVAL 573
C									
B									
Nucleotide Binding Fold II									
MOAT-D	IKDLHLVHG	CEKVGVGRT	SAGKSSMTLC	EFILSAAG	EFRLDGLMV	DGLHDLRSO	ITIPQDIL	FSGILMRL	DPGSEF 1392
MRP	IKHINVTNG	CEKVGVGRT	SAGKSSMTLC	EFILSAAG	EFRLDGLMV	DGLHDLRSO	ITIPQDIL	FSGILMRL	DPGSEF 1396
cMOAT	IRGITCDGS	CEKVGVGRT	SAGKSSMTLC	EFILSAAG	EFRLDGLMV	DGLHDLRSO	ITIPQDIL	FSGILMRL	DPGSEF 1403
MOAT-C	IKKVFETKP	CEKVGVGRT	SAGKSSMTLC	EFILSAAG	EFRLDGLMV	DGLHDLRSO	ITIPQDIL	FSGILMRL	DPGSEF 1296
MOAT-B	IKKLTALKS	CEKVGVGRT	SAGKSSMTLC	EFILSAAG	EFRLDGLMV	DGLHDLRSO	ITIPQDIL	FSGILMRL	DPGSEF 1143
SUR	IKHVNALSP	CEKVGVGRT	SAGKSSMTLC	EFILSAAG	EFRLDGLMV	DGLHDLRSO	ITIPQDIL	FSGILMRL	DPGSEF 1447
CFTR	IKHVNALSP	CEKVGVGRT	SAGKSSMTLC	EFILSAAG	EFRLDGLMV	DGLHDLRSO	ITIPQDIL	FSGILMRL	DPGSEF 1312
MDR1	IKHVNALSP	CEKVGVGRT	SAGKSSMTLC	EFILSAAG	EFRLDGLMV	DGLHDLRSO	ITIPQDIL	FSGILMRL	DPGSEF 1142
A									
MOAT-D	EDRWALSL	HLHTVSSCF	AGLDFOCSE	GENLSVGGR	LVGLARALL	SGRIIVLEA	TAALLETEN	LIC	1465
MRP	EDRWALSL	HLHTVSSCF	AGLDFOCSE	GENLSVGGR	LVGLARALL	SGRIIVLEA	TAALLETEN	LIC	1469
cMOAT	EDRWALSL	HLHTVSSCF	AGLDFOCSE	GENLSVGGR	LVGLARALL	SGRIIVLEA	TAALLETEN	LIC	1476
MOAT-C	EDRWALSL	HLHTVSSCF	AGLDFOCSE	GENLSVGGR	LVGLARALL	SGRIIVLEA	TAALLETEN	LIC	1369
MOAT-B	EDRWALSL	HLHTVSSCF	AGLDFOCSE	GENLSVGGR	LVGLARALL	SGRIIVLEA	TAALLETEN	LIC	1216
SUR	EDRWALSL	HLHTVSSCF	AGLDFOCSE	GENLSVGGR	LVGLARALL	SGRIIVLEA	TAALLETEN	LIC	1385
CFTR	EDRWALSL	HLHTVSSCF	AGLDFOCSE	GENLSVGGR	LVGLARALL	SGRIIVLEA	TAALLETEN	LIC	1520
MDR1	EDRWALSL	HLHTVSSCF	AGLDFOCSE	GENLSVGGR	LVGLARALL	SGRIIVLEA	TAALLETEN	LIC	1215
C									
B									

Fig. 6A

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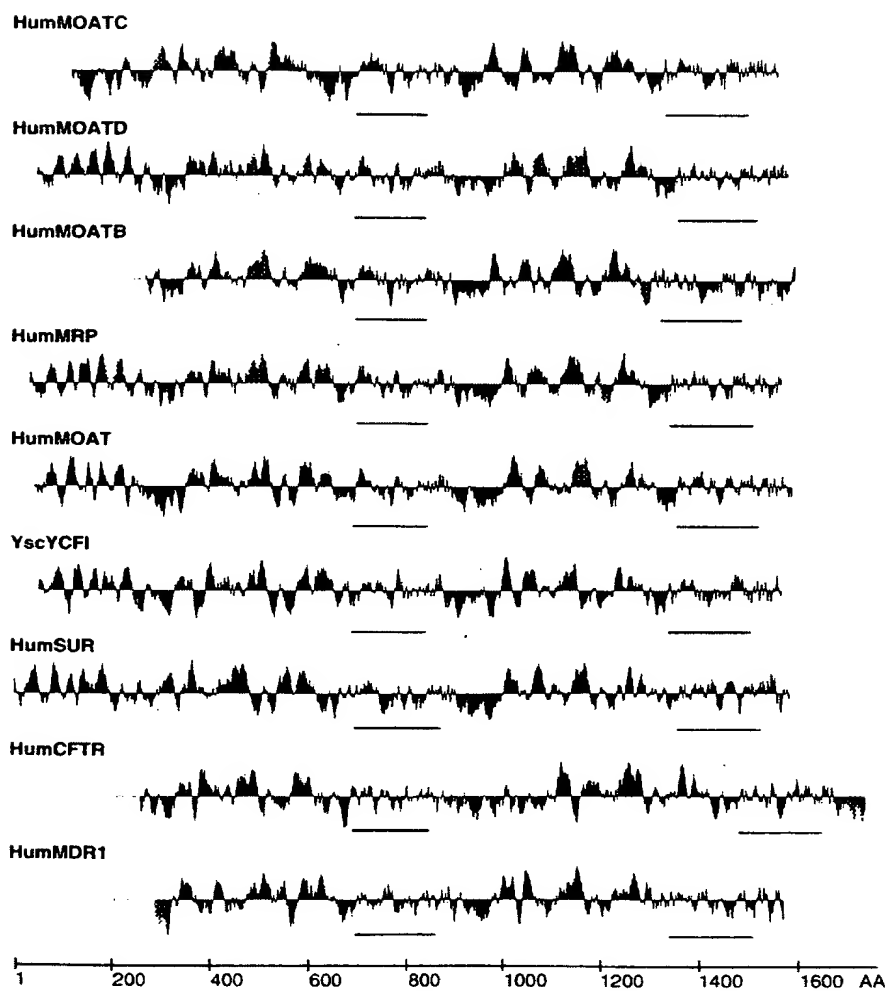


Fig. 6B

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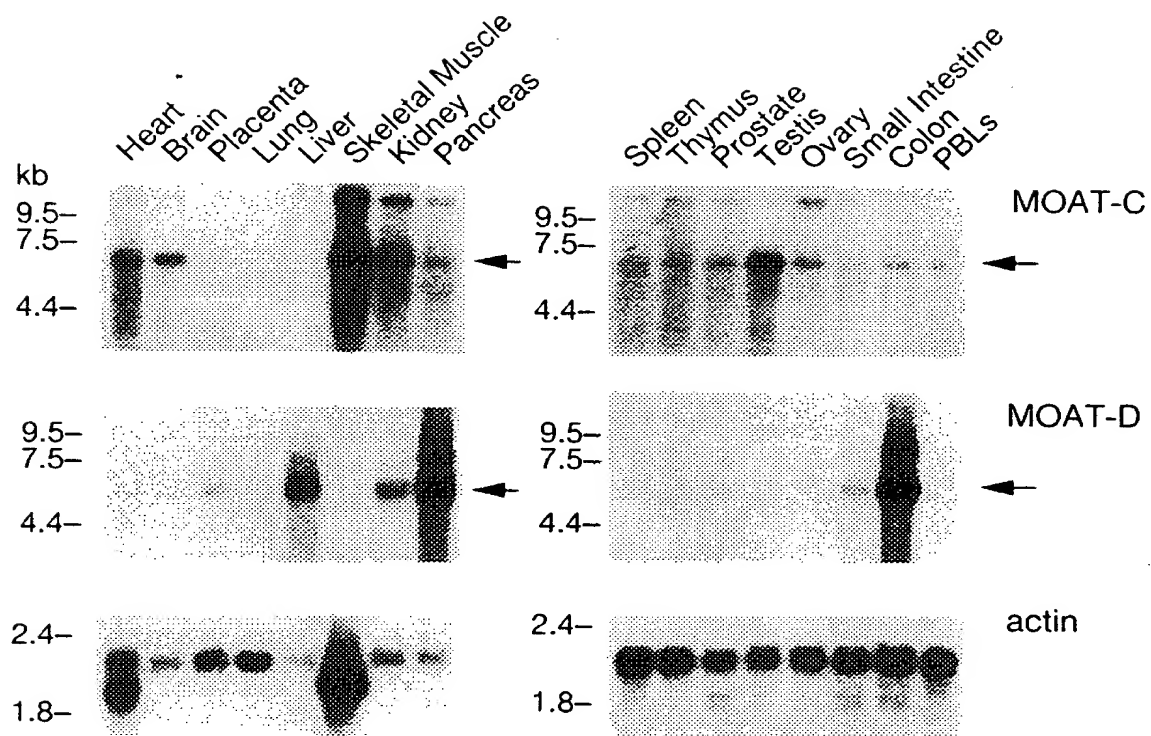


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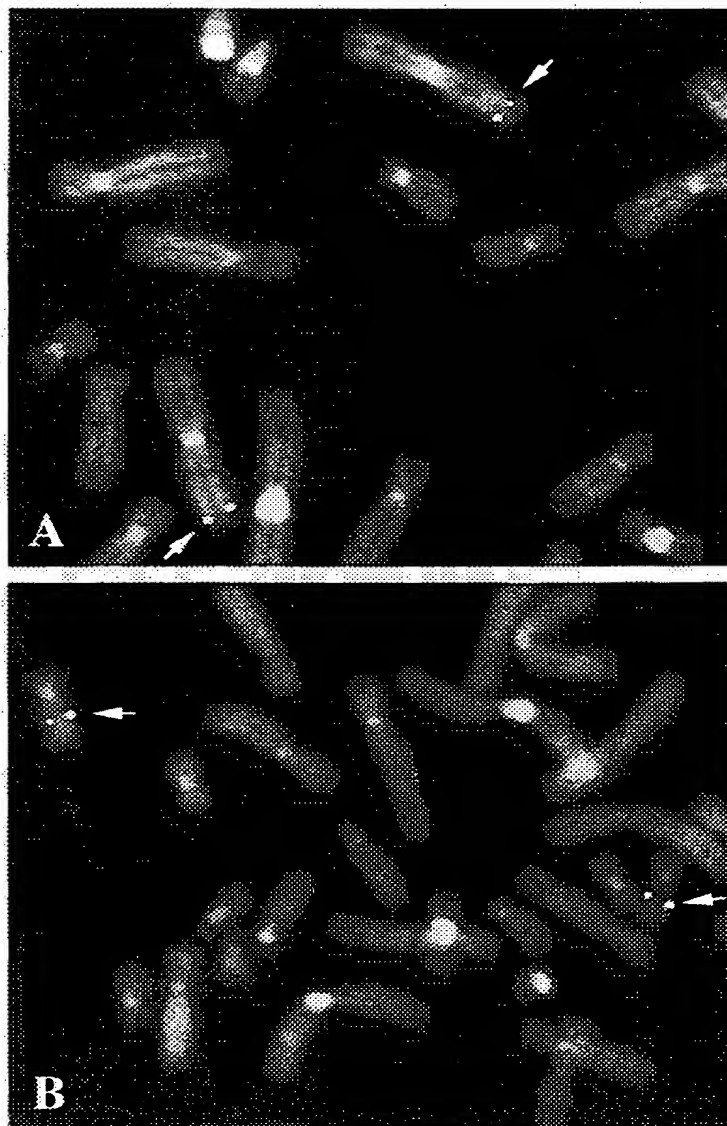


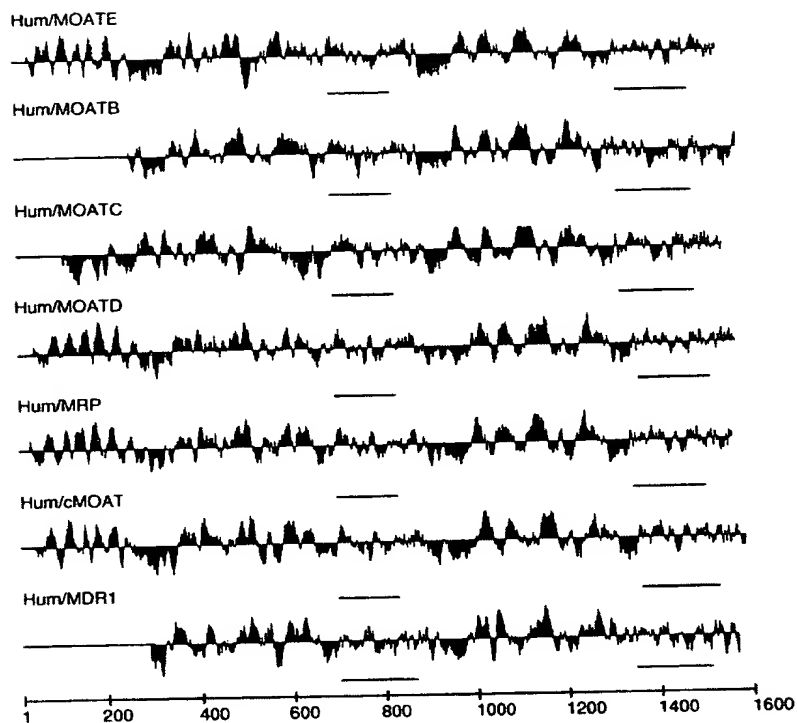
Figure 8

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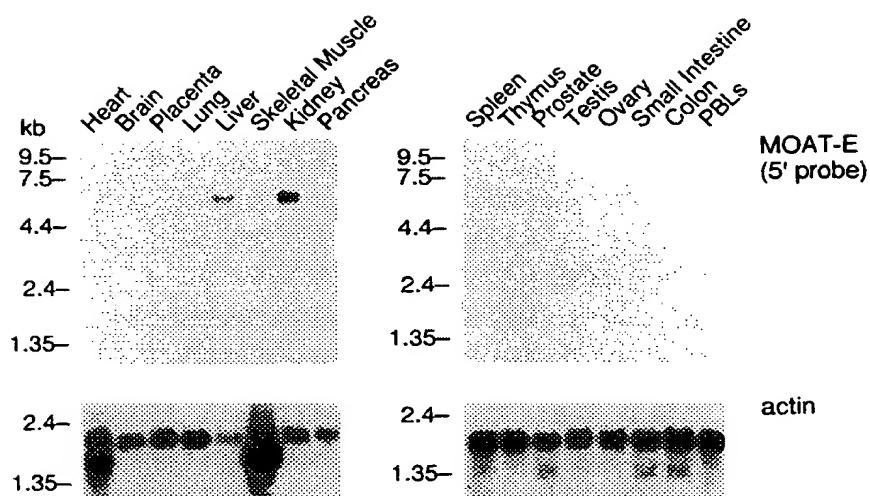
1 MAAPAEP CAG QGVWVQTEPE PAATSLLSLC FLRTAGVWVP PHYLWVLGPI YLLFIHHHGR
 61 GYLMSPLFK AKMVLGFALI VLCTSSVAVA LWKIQOGTPE APEFLIHPTV WLTTMSFAVF
 121 LIHTERKKGV QSSGVLFYVW LLCFVLPATN AAQOASGAGF QSDPVRHLST YLCLSLVVAQ
 181 FVLSC LADQP PFFPEDPQOS NPCPETGA AF PSKATFWWVS GLVWRGYRRP LRPKDLWSLG
 241 RENSSEELVS RLEKEWMNR SAARRHNKAI AFKRKGGSGM KAPETEPFLR QEGSQWRPLL
 301 KAIWQVFHST FILGTLSLII SDVFRFTVPK LLSLFLEFIG DPKPPAWKGY LLAVLMFLSA
 361 CLQTLFEQQN MYRLKVPQMR LRSATGLVY RKVLALSSGS RKASAVGDVV NLVSVDVQRL
 421 TESVLYLNGL WLPLVWIVVC FVYLWQLLGP SALTAIAVFL SLLPLNFFIS KKRNNHQEEQ
 481 MRQKDSRRL TSSILRNSKT IKPHGWEGAF LDRVLGIRGQ ELGALRTSGL LFSVSLVSFO
 541 VSTFLVALVV FAVHTLVAEN AMNAEKAFVT LTVLNILNKA QAFLPFSIHS LVQARVSFDR
 601 LVTFLCLEEV DPGVVDSSSS GSAAGKDCIT IHSATFAWSQ ESPPCLHRIN LTVPOGCLLA
 661 VVGPGVAGKS SLLSALLGEL SKVEGFVSIE GAVAYVPQEA WVQNTSVVEN VCFGQELDPP
 721 WLERVLEACA LQPDVDSFPE GIHTSIGEQG MNLSGGQKOR LSLARAVYRK AAVYLLDDPL
 781 AALDAHVGQH VFNQVIGPGG LLQGTTRILV THALHILPQA DWIIVLANGA IAEMGSYQEL
 841 LQRKGALVCL LDQARQPGDR GEGETEPGTS TKDPRGTSAG RRP ELRRERS IKSVP EKDR T
 901 TSEAQTEVPL DDPDRAGWPA GKDSIQYGRV KATVHLAYLR AVGTPLCLYA LPLFLCQOVA
 961 SFCRGYWLSL WADDPVAGGQ QTQAALRGGI FGLLGCLQAI GLFASMAAVL LGGARASRL
 1021 FQRLLDVVR SPISFFERTP IGHLLNRFSK ETDTVVDV DIP DKLRSLMYA FGLLEVSLVV
 1081 AVATPLATVA ILPLFLLYAG FQSLYVVSSC QLRRL ESASY SSVCSHMAET FQGSTVVR AF
 1141 RTQAPFVAQN NARVDESQRI SPFRLVADRW LAANVELLGN GLVF AAATCA VLSKAHLSAG
 1201 LVGFSVSAAL QVTQALQWV RNWTDLENSI VSVERMODYA WTPKEAPWRL PTCAAOPPWP
 1261 OGGQIEFRDF GLRYRPELPL AVQGVSLKIH AGEKV GIVGR TGAGKSSLAS GLLRLQEAAE
 1321 GGIWIDGVPI AHVGLHTLRS RISIIPQDPI LFPGLSRMNL DLLQEHSD EA IWA ALETVQL
 1381 KALVASLPGQ LQYKCADRGE DLSVGOKOLL CLARALLRKT OILILDEATA AVDPGTELOM
 1441 QAMLGSWFAQ CTVLLIAHRL RSVMDCARVL VMDKGQVAES GSPAQLLAQK GLFYRLAQES
 1501 GLV

Figure 9

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**Figure 10**

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**Figure 11**

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MOAT B cDNA AND AMINO ACID SEQUENCE ENCODED THEREBY

ATGCTGCCCGTGTAACAGGAGGTGAAGCCCAACCCGCTGCAGGACGCGAACATCTGCTCA
1 ----- + ----- + ----- + ----- + ----- + ----- + 60
TACGACGGGCACATGGTCCTCCACTTCGGGTGGGCGACGTCCTGCGCTTGTAGACGAGT

a M L P V Y Q E V K P N P L G D A N I C S -

CGCGTGTTCTTCTGGTGGCTCAATCCCTTGTTTAAAATTGGCCATAAACGGAGATTAGAG
61 ----- + ----- + ----- + ----- + ----- + ----- + 120
GCGCACAAGAAGACCACCGAGTTAGGGAACAAATTTTAACCGGTATTTGCCTCTAATCTC

a R V F F W W L N P L F K I G H K R R L E -

GAAGATGATATGTATTCAGTGCTGCCAGAAGACCGCTCACAGCACCTTGGAGAGGAGTTG
121 ----- + ----- + ----- + ----- + ----- + ----- + 180
CTTCTACTATACATAAGTCACGACGGTCTTCTGGCGAGTGTCGTGGAACCTCTCCTCAAC

a E D D M Y S V L P E D R S Q H L G E E L -

CAAGGGTTCTGGGATAAAGAAGTTTTAAGAGCTGAGAATGACGCACAGAAGCCTTCTTTA
181 ----- + ----- + ----- + ----- + ----- + ----- + 240
GTTCCCAAGACCCTATTTCTTCAAAATTCTCGACTCTTACTGCGTGTCTTCGGAAGAAAT

a Q G F W D K E V L R A E N D A Q K P S L -

ACAAGAGCAATCATAAAGTGTTACTGGAAATCTTATTTAGTTTTGGGAATTTTACGTTA
241 ----- + ----- + ----- + ----- + ----- + ----- + 300
TGTCTCGTTAGTATTTACAATGACCTTTAGAATAAATCAAAACCCTTAAAAATGCAAT

a T R A I I K C Y W K S Y L V L G I F T L -

ATTGAGGAAAGTGCCAAAGTAATCCAGCCCATATTTTTGGGAAAAATTATTAATTATTTT
301 ----- + ----- + ----- + ----- + ----- + ----- + 360
TAACTCCTTTCACGGTTTCATTAGGTCGGGTATAAAAACCCTTTTAAATAATTAATAAAA

a I E E S A K V I Q P I F L G K I I N Y F -

GAAAATTATGATCCCATGGATTCTGTGGCTTTGAACACAGCGTACGCCTATGCCACGGTG

Figure 12A

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361 -----+-----+-----+-----+-----+-----+ 420
 CTTTAAATACTAGGGTACCTAAGACACCGAAACTTGTGTCGCATGCGGATACGGTGCCAC

a E N Y D P M D S V A L N T A Y A Y A T V -

CTGACTTTTTGCACGCTCATTTTGGCTATACTGCATCACTTATATTTTATCACGTTTCAG

421 -----+-----+-----+-----+-----+-----+ 480
 GACTGAAAAACGTGCGAGTAAAACCGATATGACGTAGTGAATATAAAAAATAGTGCAAGTC

a L T F C T L I L A I L H H L Y F Y H V Q -

TGTGCTGGGATGAGGTTACGAGTAGCCATGTGCCATATGATTTATCGGAAGGCACTTCGT

481 -----+-----+-----+-----+-----+-----+ 540
 ACACGACCCTACTCCAATGCTCATCGGTACACGGTATACTAAATAGCCTTCCGTGAAGCA

a C A G M R L R V A M C H M I Y R K A L R -

CTTAGTAACATGGCCATGGGGAAGACAACCACAGGCCAGATAGTCAATCTGCTGTCCAAT

541 -----+-----+-----+-----+-----+-----+ 600
 GAATCATTGTACCGGTACCCCTTCTGTTGGTGTCCGGTCTATCAGTTAGACGACAGGTTA

a L S N M A M G K T T T G Q I V N L L S N -

GATGTGAACAAGTTTGATCAGGTGACAGTGTTCTTACACTTCCTGTGGGCAGGACCACTG

601 -----+-----+-----+-----+-----+-----+ 660
 CTACACTTGTTCAAAGTAGTCCACTGTCACAAGAATGTGAAGGACACCCGTCCTGGTGAC

a D V N K F D Q V T V F L H F L W A G P L -

CAGGCGATCGCAGTGACTGCCCTACTCTGGATGGAGATAGGAATATCGTGCCTTGCTGGG

661 -----+-----+-----+-----+-----+-----+ 720
 GTCCGCTAGCGTCACTGACGGGATGAGACCTACCTCTATCCTTATAGCACGGAACGACCC

a Q A I A V T A L L W M E I G I S C L A G -

ATGGCAGTTCTAATCATTCTCCTGCCCTTGCAAAGCTGTTTTGGGAAGTTGTTCTCATCA

721 -----+-----+-----+-----+-----+-----+ 780
 TACCGTCAAGATTAGTAAGAGGACGGGAACGTTTCGACAAAACCCTTCAACAAGAGTAGT

a M A V L I I L L P L Q S C F G K L F S S -

CTGAGGAGTAAAAGTCAACTTTACGGATGCCAGGATCAGGACCATGAATGAAGTTATA

781 -----+-----+-----+-----+-----+-----+ 840

Figure 12B

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GA CTCCTCATTTTGACGTTGAAAGTGCCTACGGTCCTAGTCCTGGTACTTACTTCAATAT

a L R S K T A T F T D A R I R T M N E V I -

ACTGGTATAAGGATAATAAAAAATGTACGCCTGGGAAAAGTCATTTTCAAATCTTATTACC
841 -----+-----+-----+-----+-----+-----+ 900
TGACCATATTCTATTATTTTACATGCGGACCCTTTTCAGTAAAAGTTTAGAATAATGG

a T G I R I I K M Y A W E K S F S N L I T -

AATTTGAGAAAGAAGGAGATTTC CAAGATTCTGAGAAGTTCCTGCCTCAGGGGGATGAAT
901 -----+-----+-----+-----+-----+-----+ 960
TTAAACTCTTTCTTCTCTAAAGGTTCTAAGACTCTTCAAGGACGGAGTCCCCCTACTTA

a N L R K K E I S K I L R S S C L R G M N -

TTGGCTTCGTTTTTCAGTGCAAGCAAAATCATCGTGTGTTGTGACCTTCACCACCTACGTG
961 -----+-----+-----+-----+-----+-----+ 1020
AACCGAAGCAAAAAGTCACGTTTCGTTTTAGTAGCACAAACACTGGAAGTGGTGGATGCAC

a L A S F F S A S K I I V F V T F T T Y V -

CTCCTCGGCAGTGTGATCACAGCCAGCCGCGTGTTTCGTGGCAGTGACGCTGTATGGGGCT
1021 -----+-----+-----+-----+-----+-----+ 1080
GAGGAGCCGTCACACTAGTGTCGGTCGGCGCACAAAGCACCGTCACTGCGACATACCCCGA

a L L G S V I T A S R V F V A V T L Y G A -

GTGCGGCTGACGGTTACCCTCTTCTTCCCCTCAGCCATTGAGAGGGTGTGAGAGGCAATC
1081 -----+-----+-----+-----+-----+-----+ 1140
CACGCCGACTGCCAATGGGAGAAGAAGGGGAGTCGGTAACTCTCCACAGTCTCCGTTAG

a V R L T V T L F F P S A I E R V S E A I -

GTCAGCATCCGAAGAATCCAGACCTTTTGTCTACTTGATGAGATATCACAGCGCAACCGT
1141 -----+-----+-----+-----+-----+-----+ 1200
CAGTCGTAGGCTTCTTAGGTCTGAAAAACGATGAACTACTCTATAGTGTGCGGTTGGCA

a V S I R R I Q T F L L L D E I S Q R N R -

CAGCTGCCGTCAGATGGTAAAAAGATGGTGCATGTGCAGGATTTTACTGCTTTTGGGAT
1201 -----+-----+-----+-----+-----+-----+ 1260
GTCGACGGCAGTCTACCATTTTCTACCACGTACACGTCTAAAATGACGAAAAACCCTA

Figure 12C

17/56

a Q L P S D G K K M V H V Q D F T A F W D -
AAGGCATCAGAGACCCCAACTCTACAAGGCCTTTCCTTTACTGTCAGACCTGGCGAATTG
1261 -----+-----+-----+-----+-----+ 1320
TTCCGTAGTCTCTGGGGTTGAGATGTTCCGGAAGGAAATGACAGTCTGGACCGCTTAAC

a K A S E T P T L O G L S F T V R P G E L -
TTAGCTGTGGTCGGCCCCGTGGGAGCAGGGAAGTCATCACTGTTAAGTGCCGTGCTCGGG
1321 -----+-----+-----+-----+-----+ 1380
AATCGACACCAGCCGGGGCACCCTCGTCCCTTCAGTAGTGACAATTCACGGCACGAGCCC

a L A V V G P V G A G K S S L L S A V L G -
GAATTGGCCCCAAGTCACGGGCTGGTCAGCGTGCATGGAAGAATTGCCTATGTGTCTCAG
1381 -----+-----+-----+-----+-----+ 1440
CTTAACCGGGGTTCAAGTCCCCGACCAGTCGCACGTACCTTCTTAACGGATACACAGAGTC

a E L A P S H G L V S V H G R I A Y V S Q -
CAGCCCTGGGTGTTCTCGGGAAGTCTGAGGAGTAATATTTTATTTGGGAAGAAATATGAA
1441 -----+-----+-----+-----+-----+ 1500
GTCGGGACCCACAAGAGCCCTTGAGACTCCTCATTATAAAATAAACCCCTTCTTTATACTT

a Q P W V F S G T L R S N I L F G K K Y E -
AAGGAACGATATGAAAAAGTCATAAAGGCTTGTGCTCTGAAAAAGGATTTACAGCTGTTG
1501 -----+-----+-----+-----+-----+ 1560
TTCCTTGCTATACTTTTTCAGTATTTCCGAACACGAGACTTTTTCCTAAATGTCGACAAC

a K E R Y E K V I K A C A L K K D L Q L L -
GAGGATGGTGATCTGACTGTGATAGGAGATCGGGGAACCACGCTGAGTGGAGGGCAGAAA
1561 -----+-----+-----+-----+-----+ 1620
CTCCTACCACTAGACTGACACTATCCTCTAGCCCCTTGGTGCGACTCACCTCCCGTCTTT

a E D G D L T V I G D R G T T L S G G Q K -
GCACGGGTAAACCTTGCAAGAGCAGTGTATCAAGATGCTGACATCTATCTCCTGGACGAT
1621 -----+-----+-----+-----+-----+ 1680
CGTGCCCATTTGGAACGTTCTCGTCACATAGTTCTACGACTGTAGATAGAGGACCTGCTA

Figure 12D

18/56

a A R V N L A R A V Y Q D A D I Y L L D D -
CCTCTCAGTGCAGTAGATGCGGAAGTTAGCAGACACTTGTTGGAAGTGTGTATTGTCAA
1681 -----+-----+-----+-----+-----+-----+ 1740
GGAGAGTCACGTCATCTACGCCTTCAATCGTCTGTGAACAAGCTTGACACATAAACAGTT

a P L S A V D A E V S R H L F E L C I C O -
ATTTTGCATGAGAAGATCACAATTTTAGTGACTCATCAGTTGCAGTACCTCAAAGCTGCA
1741 -----+-----+-----+-----+-----+-----+ 1800
TAAACGTACTCTTCTAGTGTTAAATCACTGAGTAGTCAACGTCATGGAGTTTCGACGT

a I L H E K I T I L V T H Q L Q Y L K A A -
AGTCAGATTCTGATATTGAAAGATGGTAAATGGTGCAGAAGGGGACTTACACTGAGTTC
1801 -----+-----+-----+-----+-----+-----+ 1860
TCAGTCTAAGACTATAACTTTCTACCATTTTACCACGTCTTCCCCTGAATGTGACTCAAG

a S Q I L I L K D G K M V Q K G T Y T E F -
CTAAAATCTGGTATAGATTTTGGCTCCCTTTTAAAGAAGGATAATGAGGAAAGTGAACAA
1861 -----+-----+-----+-----+-----+-----+ 1920
GATTTTAGACCATATCTAAAACCGAGGGAAAATTTCTTCTTACTCCTTTCACTTGT

a L K S G I D F G S L L K K D N E E S E Q -
CCTCCAGTTCAGGAACTCCACACTAAGGAATCGTACCTTCTCAGAGTCTTCGGTTTGG
1921 -----+-----+-----+-----+-----+-----+ 1980
GGAGGTCAAGGTCCTTGAGGGTGTGATTCTTAGCATGGAAGAGTCTCAGAAGCCAAACC

a P P V P G T P T L R N R T F S E S S V W -
TCTCAACAATCTTCTAGACCCTCCTTGAAAGATGGTGTCTCTGGAGAGCCAAGATACAGAG
1981 -----+-----+-----+-----+-----+-----+ 2040
AGAGTTGTTAGAAGATCTGGGAGGAACTTTCTACCACGAGACCTCTCGGTTCTATGTCTC

a S Q Q S S R P S L K D G A L E S Q D T E -
AATGTCCCAGTTACACTATCAGAGGAGAACC GTTCTGAAGGAAAAGTTGGTTTTTCAGGCC
2041 -----+-----+-----+-----+-----+-----+ 2100
TTACAGGGTCAATGTGATAGTCTCCTCTTGGAAGACTTCTTTTCAACCAAAAGTCCGG

a N V P V T L S E E N R S E G K V G F Q A

Figure 12E

SUBSTITUTE SHEET (RULE 26)

19/56

TATAAGAATTACTTCAGAGCTGGTGCTCACTGGATTGTCTTCATTTTCCTTATTCTCCTA
2101 -----+-----+-----+-----+-----+-----+ 2160
ATATTCTTAATGAAGTCTCGACCACGAGTGACCTAACAGAAGTAAAAGGAATAAGAGGAT

a Y K N Y F R A G A H W I V F I F L I L L -

AACACTGCAGCTCAGGTTGCCTATGTGCTTCAAGATTGGTGGCTTTCATACTGGGCAAAC
2161 -----+-----+-----+-----+-----+-----+ 2220
TTGTGACGTCGAGTCCAACGGATACACGAAGTTCTAACCACCGAAAGTATGACCCGTTTG

a N T A A Q V A Y V L Q D W W L S Y W A N -

AAACAAAGTATGCTAAATGTCAGTGTAAATGGAGGAGGAAATGTAACCGAGAAGCTAGAT
2221 -----+-----+-----+-----+-----+-----+ 2280
TTTGTTCATACGATTACAGTGACATTTACCTCCTTTACATTGGCTCTTCGATCTA

a K Q S M L N V T V N G G G N V T E K L D -

CTTAAGTGGTACTTAGGAATTTATTCAGGTTTAACTGTAGCTACCGTTCTTTTGGCATA
2281 -----+-----+-----+-----+-----+-----+ 2340
GAATTGACCATGAATCCTTAAATAAGTCCAAATTGACATCGATGGCAAGAAAAACCGTAT

a L N W Y L G I Y S G L T V A T V L F G I -

GCAAGATCTCTATTGGTATTCTACGTCCTTGTTAACTCTTCACAACTTTGCACAACAAA
2341 -----+-----+-----+-----+-----+-----+ 2400
CGTTCTAGAGATAACCATAAGATGCAGGAACAATTGAGAAGTGTTGAAACGTGTTGTTT

a A R S L L V F Y V L V N S S Q T L H N K -

ATGTTTGAGTCAATTCTGAAAGCTCCGGTATTATTCTTTGATAGAAATCCAATAGGAAGA
2401 -----+-----+-----+-----+-----+-----+ 2460
TACAAACTCAGTTAAGACTTTCGAGGCCATAATAAGAACTATCTTTAGGTTATCCTTCT

a M F E S I L K A P V L F F D R N P I G R -

ATTTTAAATCGTTTCTCCAAAGACATTGGACACTTGGATGATTTGCTGCCGCTGACGTTT
2461 -----+-----+-----+-----+-----+-----+ 2520
TAAAATTTAGCAAAGAGGTTTCTGTAACCTGTGAACCTACTAAACGACGGCGACTGCAAA

a I L N R F S K D I G H L D D L L P L T F

Figure 12F

20/56

TTAGATTTTCATCCAGACATTGCTACAAGTGGTTGGTGTGGTCTCTGTGGCTGTGGCCGTG
2521 -----+-----+-----+-----+-----+-----+ 2580
AATCTAAAGTAGGTCTGTAACGATGTTACCAACCACACCAGAGACACCGACACCGGCAC

a L D F I Q T L L Q V V G V V S V A V A V -

ATTCCTTGGATCGCAATACCCTTGGTTCCCTTGAATCATTTTTCATTTTCTTCGGCGA
2581 -----+-----+-----+-----+-----+-----+ 2640
TAAGGAACCTAGCGTTATGGGAACCAAGGGGAACCTTAGTAAAAGTAAAAAGAAGCCGCT

a I P W I A I P I V P L G I I F I F L R R -

TATTTTTTGGAAACGTCAAGAGATGTGAAGCGCCTGGAATCTACAACCTCGGAGTCCAGTG
2641 -----+-----+-----+-----+-----+-----+ 2700
ATAAAAAACCTTTCAGTTCTCTACACTTCGCGGACCTTAGATGTTGAGCCTCAGGTCAC

a Y F L E T S R D V K R L E S T T R S P V -

TTTTCCCACTTGTCATCTTCTCTCCAGGGGCTCTGGACCATCCGGGCATACAAAGCAGAA
2701 -----+-----+-----+-----+-----+-----+ 2760
AAAAGGGTGAACAGTAGAAGAGAGGTCCCCGAGACCTGGTAGGCCCGTATGTTTCGTCTT

a F S H L S S S L Q G L W T I R A Y K A E -

GAGAGGTGTCAGGAAGTGTGATGCACACCAGGATTTACATTCAGAGGCTTGGTTCTTG
2761 -----+-----+-----+-----+-----+-----+ 2820
CTCTCCACAGTCCTTGACAAACTACGTGTGGTCTAAATGTAAGTCTCCGAACCAAGAAC

a E R C Q E L F D A H Q D L H S E A W F L -

TTTTTGACAACGTCCCGCTGGTTCGCCGTCCGTCTGGATGCCATCTGTGCCATGTTTGTC
2821 -----+-----+-----+-----+-----+-----+ 2880
AAAAACTGTTGCAGGGCGACCAAGCGGCAGGCAGACCTACGGTAGACACGGTACAAACAG

a F L T T S R W F A V R L D A I C A M F V -

ATCATCGTTGCCTTTGGGTCCCTGATTCTGGCAAAAACCTCTGGATGCCGGGCAGGTTGGT
2881 -----+-----+-----+-----+-----+-----+ 2940
TAGTAGCAACGGAAACCCAGGGACTAAGACCGTTTTTGAGACCTACGGCCCGTCCAACCA

a I I V A F G S L I L A K T L D A G Q V G -

TTGGCACTGTCCTATGCCCTCACGCTCATGGGGATGTTTCAGTGGTGTGTTTCGACAAAGT

Figure 12G

SUBSTITUTE SHEET (RULE 26)

21/56

2941 -----+-----+-----+-----+-----+-----+ 3000
AACCGTGACAGGATACGGGAGTGCGAGTACCCCTACAAAGTCACCACACAAGCTGTTTCA

a L A L S Y A L T L M G M F Q W C V R Q S .

GCTGAAGTTGAGAATATGATGATCTCAGTAGAAAGGGTCATTGAATACACAGACCTTGAA

3001 -----+-----+-----+-----+-----+-----+ 3060
CGACTTCAACTCTTATACTACTAGAGTCATCTTTCCCAGTAACTTATGTGTCTGGAACCT

a A E V E N M M I S V E R V I E Y T D L E .

AAAGAAGCACCTTGGAATATCAGAAACGCCCACCACCAGCCTGGCCCCATGAAGGAGTG

3061 -----+-----+-----+-----+-----+-----+ 3120
TTTCTTCGTGGAACCTTATAGTCTTTGCGGGTGGTGGTCGGACCGGGTACTTCCTCAC

a K E A P W E Y Q K R P P P A W P H E G V .

ATAATCTTTGACAATGTGAACTTCATGTACAGTCCAGGTGGGCCTCTGGTACTGAAGCAT

3121 -----+-----+-----+-----+-----+-----+ 3180
TATTAGAACTGTTACACTTGAAGTACATGTCAGGTCCACCCGGAGACCATGACTTCGTA

a I I F D N V N F M Y S P G G P L V L K H .

CTGACAGCACTCATTAAATCACAAGAAAAGGTTGGCATTGTGGGAAGAACCGGAGCTGGA

3181 -----+-----+-----+-----+-----+-----+ 3240
GACTGTCGTGAGTAATTTAGTGTCTTTTCCAACCGTAACACCCTTCTTGGCCTCGACCT

a L T A L I K S Q E K V G I V G R T G A G .

AAAAGTTCCCTCATCTCAGCCCTTTTTAGATTGTCAGAACCCGAAGGTAAAATTTGGATT

3241 -----+-----+-----+-----+-----+-----+ 3300
TTTTCAAGGGAGTAGAGTCGGGAAAAATCTAACAGTCTTGGGCTTCCATTTTAAACCTAA

a K S S L I S A L F R L S E P E G K I W I .

GATAAGATCTTGACAACTGAAATTGGACTTCACGATTTAAGGAAGAAAATGTCAATCATA

3301 -----+-----+-----+-----+-----+-----+ 3360
CTATTCTAGAACTGTTGACTTTAACCTGAAGTGCTAAATTCCTTCTTTTACAGTTAGTAT

a D K I L T T E I G L H D L R K K M S I I .

CCTCAGGAACCTGTTTTGTTCACTGGAACAATGAGGAAAAACCTGGATCCCTTTAAGGAG

3361 -----+-----+-----+-----+-----+-----+ 3420

Figure 12H

22/56

GGAGTCCTTGGACAAAACAAGTGACCTTGTTACTCCTTTTTGGACCTAGGGAAATTCCTC

a P Q E P V L F T G T M R K N L D P F K E -

CACACGGATGAGGAACTGTGGAATGCCTTACAAGAGGTACAACCTAAAGAAACCATTGAA

3421 -----+-----+-----+-----+-----+ 3480

GTGTGCCTACTCCTTGACACCTTACGGAATGTTCTCCATGTTGAATTTCTTTGGTAACTT

a H T D E E L W N A L Q E V Q L K E T I E -

GATCTTCCTGGTAAAATGGATACTGAATTAGCAGAATCAGGATCCAATTTTAGTGTTGGA

3481 -----+-----+-----+-----+-----+ 3540

CTAGAAGGACCATTTTACCTATGACTTAATCGTCTTAGTCCTAGGTAAAATCACAACT

a D L P G K M D T E L A E S G S N F S V G -

CAAAGACAACTGGTGTGCCTTGCCAGGGCAATTCTCAGGAAAAATCAGATATTGATTATT

3541 -----+-----+-----+-----+-----+ 3600

GTTTCTGTTGACCACACGGAACGGTCCCGTTAAGAGTCCTTTTTAGTCTATAACTAATAA

a Q R Q L V C L A R A I L R K N Q I L I I -

GATGAAGCGACGGCAAATGTGGATCCAAGAACTGATGAGTTAATACAAAAAAAAATCCGG

3601 -----+-----+-----+-----+-----+ 3660

CTACTTCGCTGCCGTTTACACCTAGGTTCTTGACTACTCAATTATGTTTTTTTTTAGGCC

a D E A T A N V D P R T D E L I Q K K I R -

GAGAAATTTGCCCACTGCACCGTGCTAACCATTGCACACAGATTGAACACCATTATTGAC

3661 -----+-----+-----+-----+-----+ 3720

CTCTTTAAACGGGTGACGTGGCAGATTGGTAACGTGTGTCTAACTTGTTGGTAATAACTG

a E K F A H C T V L T I A H R L N T I I D -

AGCGACAAGATAATGGTTTTAGATTGAGGAAGACTGAAAGAATATGATGAGCCGTATGTT

3721 -----+-----+-----+-----+-----+ 3780

TCGCTGTTCTATTACCAAATCTAAGTCCTTCTGACTTTCTTATACTACTCGGCATACAA

a S D K I M V L D S G R L K E Y D E P Y V -

TTGCTGCAAAATAAAGAGAGCCTATTTTACAAGATGGTGCAACAACTGGGCAAGGCAGAA

3781 -----+-----+-----+-----+-----+ 3840

AACGACGTTTTATTTCTCTCGGATAAAATGTTCTACCACGTTGTTGACCCGTTCCGTCTT

Figure 12I

SUBSTITUTE SHEET (RULE 26)

23/56

a L L Q N K E S L F Y K M V Q Q L G K A E -

GCCGCTGCCCTCACTGAAACAGCAAAACAGGTATACTTCAAAAGAAATTATCCACATATT
3841 -----+-----+-----+-----+-----+-----+ 3900
CGGCGACGGGAGTGACTTTGTCGTTTTGTCCATATGAAGTTTTCTTTAATAGGTGTATAA

a A A A L T E T A K Q V Y F K R N Y P H I -

GGTCACACTGACCACATGGTTACAAACACTTCCAATGGACAGCCCTCGACCTTAACTATT
3901 -----+-----+-----+-----+-----+-----+ 3960
CCAGTGTGACTGGTGTACCAATGTTTGTGAAGGTTACCTGTCGGGAGCTGGAATTGATAA

a G H T D H M V T N T S N G Q P S T L T I -

TTCGAGACAGCACTG
3961 -----+----- 3975
AAGCTCTGTCGTGAC

a F E T A L -

Figure 12J

24/56

MOAT C cDNA AND AMINO ACID SEQUENCE ENCODED THEREBY

ATGAAGGATATCGACATAGGAAAAGAGTATATCATCCCCAGTCCTGGGTATAGAAGTGTG
1 -----+-----+-----+-----+-----+-----+ 60
TACTTCCTATAGCTGTATCCTTTTCTCATATAGTAGGGGTCAGGACCCATATCTTCACAC

a M K D I D I G K E Y I I P S P G Y R S V -

AGGGAGAGAACCAGCACTTCTGGGACGCACAGAGACCGTGAAGATTCCAAGTTCAGGAGA
61 -----+-----+-----+-----+-----+-----+ 120
TCCCTCTCTTGGTCGTGAAGACCCTGCGTGTCTCTGGCACTTCTAAGGTTCAAGTCCTCT

a R E R T S T S G T H R D R E D S K F R R -

ACTCGACCGTTGGAATGCCAAGATGCCTTGGAAACAGCAGCCCGAGCCGAGGGCCTCTCT
121 -----+-----+-----+-----+-----+-----+ 180
TGAGCTGGCAACCTTACGGTTCTACGGAACCTTTGTCGTCGGGCTCGGCTCCCGGAGAGA

a T R P L E C Q D A L E T A A R A E G L S -

CTTGATGCCTCCATGCATTCTCAGCTCAGAATCCTGGATGAGGAGCATCCCAAGGGAAAG
181 -----+-----+-----+-----+-----+-----+ 240
GAACTACGGAGGTACGTAAGAGTCGAGTCTTAGGACCTACTCCTCGTAGGGTTCCCTTTC

a L D A S M H S Q L R I L D E E H P K G K -

TACCATCATGGCTTGAGTGCTCTGAAGCCCATCCGGACTACTTCCAAACACCAGCACCCA
241 -----+-----+-----+-----+-----+-----+ 300
ATGGTAGTACCGAACTCACGAGACTTCGGGTAGGCCTGATGAAGGTTTGTGGTCGTGGGT

a Y H H G L S A L K P I R T T S K H Q H P -

GTGGACAATGCTGGGCTTTTTTCTGTATGACTTTTTCTGGGCTTTCTTCTCTGGCCCGT
301 -----+-----+-----+-----+-----+-----+ 360
CACCTGTTACGACCCGAAAAAAGGACATACTGAAAAAGCACCGAAAGAAGAGACCGGGCA

a V D N A G L F S C M T F S W L S S L A R -

GTGGCCCACAAGAAGGGGGAGCTCTCAATGGAAGACGTGTGGTCTCTGTCCAAGCACGAG

Figure 13A

25/56

361 -----+-----+-----+-----+-----+-----+ 420
CACCGGGTGTCTTCCCCCTCGAGAGTTACCTTCTGCACACCAGAGACAGGTTTCGTGCTC

a V A H K K G E L S M E D V W S L S K H E .

TCTTCTGACGTGAACTGCAGAAGACTAGAGAGACTGTGGCAAGAAGAGCTGAATGAAGTT

421 -----+-----+-----+-----+-----+-----+ 480
AGAAGACTGCACTTGACGTCTTCTGATCTCTCTGACACCGTTCTTCTCGACTTACTTCAA

a S S D V N C R R L E R L W Q E E L N E V .

GGGCCAGACGCTGCTTCCCTGCGAAGGGTTGTGTGGATCTTCTGCCGCACCAGGCTCATC

481 -----+-----+-----+-----+-----+-----+ 540
CCCGGTCTGCGACGAAGGGACGTTCCCAACACACCTAGAAGACGGCGTGGTCCGAGTAG

a G P D A A S L R R V V W I F C R T R L I .

CTGTCCATCGTGTGCCTGATGATCACGCAGCTGGCTGGCTTCAGTGGACCAGCCTTCATG

541 -----+-----+-----+-----+-----+-----+ 600
GACAGGTAGCACACGGACTACTAGTGCCTCGACCGACCGAAGTCACCTGGTCGGAAGTAC

a L S I V C L M I T Q L A G F S G P A F M .

GTGAAACACCTCTTGGAGTATACCCAGGCAACAGAGTCTAACCTGCAGTACAGCTTGTTG

601 -----+-----+-----+-----+-----+-----+ 660
CACTTTGTGGAGAACCTCATATGGGTCCGTTGTCTCAGATTGGACGTCATGTGGAACAAC

a V K H L L E Y T Q A T E S N L Q Y S L L .

TTAGTGCTGGGCCTCCTCCTGACGGAAATCGTGCGGTCTTGGTCGCTTGCACTGACTTGG

661 -----+-----+-----+-----+-----+-----+ 720
AATCACGACCCGGAGGAGGACTGCCTTTAGCACGCCAGAACCAGCGAACGTGACTGAACC

a L V L G L L L T E I V R S W S L A L T W .

GCATTGAATTACCGAACCAGGTGTCGCTTGCGGGGGGCCATCCTAACCATGGCATTTAAG

721 -----+-----+-----+-----+-----+-----+ 780
CGTAACTTAATGGCTTGGCCACAGGCGAACGCCCCCGGTAGGATTGGTACCGTAAATTC

a A L N Y R T G V R L R G A I L T M A F K .

AAGATCCTTAAGTTAAAGAACATTAAGAGAAATCCCTGGGTGAGCTCATCAACATTTGC

781 -----+-----+-----+-----+-----+-----+ 840

Figure 13B

SUBSTITUTE SHEET (RULE 26)

26/56

TTCTAGGAATTCAATTTCTTGTAATTTCTCTTTAGGGACCCACTCGAGTAGTTGTAAACG

a K I L K L K N I K E K S L G E L I N I C -

TCCAACGATGGGCAGAGAATGTTTGAGGCAGCAGCCGTTGGCAGCCTGCTGGCTGGAGGA
 841 -----+-----+-----+-----+-----+-----+ 900
 AGGTTGCTACCCGCTCTCTTACAAACTCCGTCGTCGCAACCGTCGGACGACCGACCTCCT

a S N D G Q R M F E A A A V G S L L A G G -

CCCGTTGTTGCCATCTTAGGCATGATTATAATGTAATTATTCTGGGACCAACAGGCTTC
 901 -----+-----+-----+-----+-----+-----+ 960
 GGGCAACAACGGTAGAATCCGTAATAATATTACATTAATAAGACCCTGGTTGTCCGAAG

a P V V A I L G M I Y N V I I L G P T G F -

CTGGGATCAGCTGTTTTATCCTCTTTTACCCAGCAATGATGTTTGCATCACGGCTCACA
 961 -----+-----+-----+-----+-----+-----+ 1020
 GACCCTAGTCGACAAAAATAGGAGAAAATGGGTCGTTACTACAAACGTAGTGCCGAGTGT

a L G S A V F I L F Y P A M M F A S R L T -

GCATATTTACAGGAGAAAATGCGTGGCCGCCACGGATGAACGTGTCCAGAAGATGAATGAA
 1021 -----+-----+-----+-----+-----+-----+ 1080
 CGTATAAAGTCCTCTTTTACGCACCGGCGGTGCCTACTTGACACAGGTCTTCTACTTACTT

a A Y F R R K C V A A T D E R V Q K M N E -

GTTCTTACTTACATTAAATTTATCAAAATGTATGCCTGGGTCAAAGCATTTTCTCAGAGT
 1081 -----+-----+-----+-----+-----+-----+ 1140
 CAAGAATGAATGTAATTTAAATAGTTTTACATACGGACCCAGTTTCGTAAAAGAGTCTCA

a V L T Y I K F I K M Y A W V K A F S Q S -

GTTTCAGAAAATCCGCGAGGAGGAGCGTCGGATATTGGAAAAAGCCGGGTACTTCCAGGGT
 1141 -----+-----+-----+-----+-----+-----+ 1200
 CAAGTCTTTTAGGCGCTCCTCCTCGCAGCCTATAACCTTTTTTCGGCCCATGAAGGTCCCA

a V Q K I R E E E R R I L E K A G Y F Q G -

ATCACTGTGGGTGTGGCTCCCATTTGTGGTGGTGATTGCCAGCGTGGTGACCTTCTCTGTT
 1201 -----+-----+-----+-----+-----+-----+ 1260
 TAGTGACACCCACACCGAGGGTAACACCACCACTAACGGTCGCACCACTGGAAGAGACAA

Figure 13C

SUBSTITUTE SHEET (RULE 26)

27/56

a I T V G V A P I V V V I A S V V T F S V .
CATATGACCCTGGGCTTCGATCTGACAGCAGCACAGGCTTTTACAGTGGTGACAGTCTTC
1261 ----- + ----- + ----- + ----- + ----- + ----- + 1320
GTATACTGGGACCCGAAGCTAGACTGTCGTCGTGTCCGAAAGTGTCACTACTGTCAGAAG

a H M T L G F D L T A A Q A F T V V T V F .
AATCCATGACTTTTGCTTTGAAAGTAACACCGTTTTTTCAGTAAAGTCCCTCTCAGAAGCC
1321 ----- + ----- + ----- + ----- + ----- + ----- + 1380
TTAAGGTACTGAAAACGAAACTTTTCATTGTGGCAAAAGTCATTTTCAGGGAGAGTCTTCGG

a N S M T F A L K V T P F S V K S L S E A .
TCAGTGGCTGTTGACAGATTTAAGAGTTTGTCTTAATGGAAGAGGTTACATGATAAAG
1381 ----- + ----- + ----- + ----- + ----- + ----- + 1440
AGTCACCGACAACCTGTCTAAATTCTCAAACAAAGATTACCTTCTCCAAGTGTACTATTTTC

a S V A V D R F K S L F L M E E V H M I K .
AACAAACCAGCCAGTCCTCACATCAAGATAGAGATGAAAAATGCCACCTTGGCATGGGAC
1441 ----- + ----- + ----- + ----- + ----- + ----- + 1500
TTGTTTGGTCGGTCAGGAGTGTAGTTCTATCTCTACTTTTTACGGTGGAACCGTACCCTG

a N K P A S P H I K I E M K N A T L A W D .
TCCTCCCACTCCAGTATCCAGAACTCGCCCAAGCTGACCCCCAAAAATGAAAAAAGACAAG
1501 ----- + ----- + ----- + ----- + ----- + ----- + 1560
AGGAGGGTGAGGTCATAGGTCTTGAGCGGGTTCGACTGGGGGTTTTACTTTTTTCTGTTC

a S S H S S I Q N S P K L T P K M K K D K .
AGGGCTTCCAGGGGCAAGAAAGAGAAGGTGAGGCAGCTGCAGCGCACTGAGCATCAGGCG
1561 ----- + ----- + ----- + ----- + ----- + ----- + 1620
TCCCGAAGGTCCCCGTTCTTTCTCTTCCACTCCGTCGACGTCGCGTGACTCGTAGTCCGC

a R A S R G K K E K V R Q L Q R T E H Q A .
GTGCTGGCAGAGCAGAAAGGCCACCTCCTCTGGACAGTGACGAGCGGCCAGTCCCGAA
1621 ----- + ----- + ----- + ----- + ----- + ----- + 1680
CACGACCGTCTCGTCTTTCCGGTGGAGGAGGACCTGTCACTGCTCGCCGGGTCAGGGCTT

Figure 13D

SUBSTITUTE SHEET (RULE 26)

28/56

a V L A E Q K G H L L L D S D E R P S P E -
GAGGAAGAAGGCAAGCACATCCACCTGGGCCACCTGCGCTTACAGAGGACACTGCACAGC
1681 -----+-----+-----+-----+-----+-----+ 1740
CTCCTTCTTCCGTTTCGTGTAGGTGGACCCGGTGGACGCGAATGTCTCCTGTGACGTGTGCG

a E E E G K H I H L G H L R L O R T L H S -
ATCGATCTGGAGATCCAAGAGGGTAAACTGGTTGGAATCTGCGGCAGTGTGGGAAGTGGGA
1741 -----+-----+-----+-----+-----+-----+ 1800
TAGCTAGACCTCTAGGTTCTCCCATTTGACCAACCTTAGACGCCGTCACACCCTTCACCT

a I D L E I Q E G K L V G I C G S V G S G -
AAACCTCTCTCATTTAGCCATTTTAGGCCAGATGACGCTTCTAGAGGGCAGCATTGCA
1801 -----+-----+-----+-----+-----+-----+ 1860
TTTTGGAGAGAGTAAAGTCGGTAAATCCGGTCTACTGCGAAGATCTCCCGTCGTAACGT

a K T S L I S A I L G Q M T L L E G S I A -
ATCAGTGGAAACCTTCGCTTATGTGGCCCAGCAGGCCTGGATCCTCAATGCTACTCTGAGA
1861 -----+-----+-----+-----+-----+-----+ 1920
TAGTCACCTTGGGAAGCGAATACACCGGGTTCGTCGGACCTAGGAGTTACGATGAGACTCT

a I S G T F A Y V A Q Q A W I L N A T L R -
GACAACATCCTGTTTGGGAAGGAATATGATGAAGAAAGATACAACTCTGTGCTGAACAGC
1921 -----+-----+-----+-----+-----+-----+ 1980
CTGTTGTAGGACAAACCCTTCCTTATACTACTTCTTTCTATGTTGAGACACGACTTGTGCG

a D N I L F G K E Y D E E R Y N S V L N S -
TGCTGCCTGAGGCCTGACCTGGCCATTCTTCCCAGCAGCGACCTGACGGAGATTGGAGAG
1981 -----+-----+-----+-----+-----+-----+ 2040
ACGACGGACTCCGGACTGGACCGGTAAGAAGGGTCGTCGCTGGACTGCCTCTAACCTCTC

a C C L R P D L A I L P S S D L T E I G E -
CGAGGAGCCAACCTGAGCGGTGGGCAGCGCCAGAGGATCAGCCTTGCCCCGGGCCTTGAT
2041 -----+-----+-----+-----+-----+-----+ 2100
GCTCCTCGGTTGGACTCGCCACCCGTCGCGGTCTCCTAGTCGGAACGGGCCCGGAACATA

a R G A N L S G G O R O R I S L A R A L Y -

Figure 13E

29/56

AGTGACAGGAGCATCTACATCCTGGACGACCCCTCAGTGCCTTAGATGCCCATGTGGGC
2101 -----+-----+-----+-----+-----+ 2160
TCACTGTCCTCGTAGATGTAGGACCTGCTGGGGGAGTCACGGAATCTACGGGTACACCCG

a S D R S I Y I L D D P L S A L D A H V G -

AACCACATCTTCAATAGTGCTATCCGGAACATCTCAAGTCCAAGACAGTTCTGTTTGT
2161 -----+-----+-----+-----+-----+ 2220
TTGGTGTAGAAGTTATCACGATAGGCCTTTGTAGAGTTCAGGTTCTGTCAAGACAAACAA

a N H I F N S A I R K H L K S K T V L F V -

ACCCACCAGTTACAGTACCTGGTTGACTGTGATGAAGTGATCTTCATGAAAGAGGGCTGT
2221 -----+-----+-----+-----+-----+ 2280
TGGGTGGTCAATGTCATGGACCAACTGACACTACTTCACTAGAAGTACTTTCTCCCGACA

a T H Q L Q Y L V D C D E V I F M K E G C -

ATTACGGAAAGAGGCACCCATGAGGAACTGATGAATTTAAATGGTGACTATGCTACCATT
2281 -----+-----+-----+-----+-----+ 2340
TAATGCCTTTCTCCGTGGGTACTCCTTGACTACTTAAATTTACCACTGATACGATGGTAA

a I T E R G T H E E L M N L N G D Y A T I -

TTTAATAACCTGTTGCTGGGAGAGACACCGCCAGTTGAGATCAATTCAAAAAAGGAAACC
2341 -----+-----+-----+-----+-----+ 2400
AAATTATTGGACAACGACCCTCTCTGTGGCGGTCAACTCTAGTTAAGTTTTTCTTTGG

a F N N L L L G E T P P V E I N S K K E T -

AGTGGTTCACAGAAGAAGTCACAAGACAAGGGTCCTAAAACAGGATCAGTAAAGAAGGAA
2401 -----+-----+-----+-----+-----+ 2460
TCACCAAGTGTCTTCTTCAGTGTTCTGTTCCAGGATTTGTCTTAGTCATTTCTTCTT

a S G S Q K K S Q D K G P K T G S V K K E -

AAAGCAGTAAAGCCAGAGGAAGGGCAGCTTGTGCAGCTGGAAGAGAAAGGGCAGGGTTCA
2461 -----+-----+-----+-----+-----+ 2520
TTTCGTCAATTCGGTCTCCTTCCCGTCGAACACGTCGACCTTCTTTCCCGTCCCAAGT

a K A V K P E E G Q L V Q L E E K G Q G S -

Figure 13F

SUBSTITUTE SHEET (RULE 26)

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GTGCCCTGGTCAGTATATGGTGTCTACATCCAGGCTGCTGGGGGGCCCCTTGGCATTCCCTG
2521 -----+-----+-----+-----+-----+ 2580
CACGGGACCAGTCATATACCACAGATGTAGGTCCGACGACCCCCGGGGAACCGTAAGGAC

a V P W S V Y G V Y I Q A A G G P L A F L .

GTTATTATGGCCCTTTTCATGCTGAATGTAGGCAGCACCGCCTTCAGCACCTGGTGGTTG
2581 -----+-----+-----+-----+-----+ 2640
CAATAATACCGGGAAGTACGACTTACATCCGTCGTGGCGGAAGTCGTGGACCACCAAC

a V I M A L F M L N V G S T A F S T W W L .

AGTTACTGGATCAAGCAAGGAAGCGGGAACACCACTGTGACTCGAGGGAACGAGACCTCG
2641 -----+-----+-----+-----+-----+ 2700
TCAATGACCTAGTTCGTTCCCTTCGCCCTTGTGGTGACACTGAGCTCCCTTGCTCTGGAGC

a S Y W I K Q G S G N T T V T R G N E T S .

GTGAGTGACAGCATGAAGGACAATCCTCATATGCAGTACTATGCCAGCATCTACGCCCTC
2701 -----+-----+-----+-----+-----+ 2760
CACTCACTGTCGTA CTTCCTGTTAGGAGTATACGTCATGATACGGTCGTAGATGCGGGAG

a V S D S M K D N P H M Q Y Y A S I Y A L .

TCCATGGCAGTCATGCTGATCCTGAAAGCCATTCGAGGAGTTGTCTTTGTCAAGGGCACG
2761 -----+-----+-----+-----+-----+ 2820
AGGTACCGTCAGTACGACTAGGACTTTCGGTAAGCTCCTCAACAGAAACAGTTCCCGTGC

a S M A V M L I L K A I R G V V F V K G T .

CTGCGAGCTTCCTCCCGGCTGCATGACGAGCTTTTCCGAAGGATCCTTCGAAGCCCTATG
2821 -----+-----+-----+-----+-----+ 2880
GACGCTCGAAGGAGGGCCGACGTA CTGCTCGAAAAGGCTTCCTAGGAAGCTTCGGGATAC

a L R A S S R L H D E L F R R I L R S P M .

AAGTTTTTTGACACGACCCCCACAGGGAGGATTCTCAACAGGTTTTCCAAAGACATGGAT
2881 -----+-----+-----+-----+-----+ 2940
TTCAAAAACTGTGCTGGGGGTGTCCCTCCTAAGAGTTGTCCAAAAGGTTTCTGTACCTA

a K F F D T T P T G R I L N R F S K D M D .

GAAGTTGACGTGCGGCTGCCGTTCCAGGCCGAGATGTTTCATCCAGAACGTTATCCTGGTG

Figure 13G

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2941 -----+-----+-----+-----+-----+-----+ 3000
 CTTCAACTGCACGCCGACGGCAAGGTCCGGCTCTACAAGTAGGTCTTGCAATAGGACCAC

a E V D V R L P F Q A E M F I Q N V I L V -

TTCTTCTGTGTGGGAATGATCGCAGGAGTCTTCCCGTGGTTCCTTGTGGCAGTGGGGCCC
 3001 -----+-----+-----+-----+-----+-----+ 3060
 AAGAAGACACACCCTTACTAGCGTCCTCAGAAGGGCACCAAGGAACACCGTCACCCCGGG

a F F C V G M I A G V F P W F L V A V G P -

CTTGTATCCTCTTTTCAGTCCTGCACATTGTCTCCAGGGTCTGATTCGGGAGCTGAAG
 3061 -----+-----+-----+-----+-----+-----+ 3120
 GAACAGTAGGAGAAAAGTCAGGACGTGTAACAGAGGTCCCAGGACTAAGCCCTCGACTTC

a L V I L F S V L H I V S R V L I R E L K -

CGTCTGGACAATATCACGCAGTCACCTTTCTCTCCACATCACGTCCAGCATAACAGGGC
 3121 -----+-----+-----+-----+-----+-----+ 3180
 GCAGACCTGTTATAGTGCGTCAGTGGAAGGAGAGGGTGTAGTGCAGGTCGTATGTCCCG

a R L D N I T Q S P F L S H I T S S I Q G -

CTTGCCACCATCCACGCCTACAATAAAGGGCAGGAGTTTCTGCACAGATACCAGGAGCTG
 3181 -----+-----+-----+-----+-----+-----+ 3240
 GAACGGTGGTAGGTGCGGATGTTATTTCCCGTCTCAAAGACGTGTCTATGGTCCTCGAC

a L A T I H A Y N K G Q E F L H R Y Q E L -

CTGGATGACAACCAAGCTCCTTTTTTTTTTTGTTTACGTGTGCGATGCGGTGGCTGGCTGTG
 3241 -----+-----+-----+-----+-----+-----+ 3300
 GACCTACTGTTGGTTCGAGGAAAAAAAAACAAATGCACACGCTACGCCACCGACCGACAC

a L D D N Q A P F F L F T C A M R W L A V -

CGGCTGGACCTCATCAGCATCGCCCTCATCACCACCACGGGGCTGATGATCGTTCTTATG
 3301 -----+-----+-----+-----+-----+-----+ 3360
 GCCGACCTGGAGTAGTCGTAGCGGGAGTAGTGGTGGTGCCCCGACTACTAGCAAGAATAC

a R L D L I S I A L I T T T G L M I V L M -

CACGGGCAGATTCCCCCAGCCTATGCGGGTCTCGCCATCTCTTATGCTGTCCAGTTAAGC
 3361 -----+-----+-----+-----+-----+-----+ 3420

Figure 13H

SUBSTITUTE SHEET (RULE 26)

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GTGCCCCGTCTAAGGGGGTCGGATACGCCAGAGCGGTAGAGAATACGACAGGTCAATTGC

a H G Q I P P A Y A G L A I S Y A V Q L T -

GGGCTGTTCCAGTTTACGGTCAGACTGGCATCTGAGACAGAAGCTCGATTACCTCGGTG
3421 -----+-----+-----+-----+-----+-----+ 3480
CCCGACAAGGTCAAATGCCAGTCTGACCGTAGACTCTGTCTTCGAGCTAAGTGGAGCCAC

a G L F O F T V R L A S E T E A R F T S V -

GAGAGGATCAATCACTACATTAAGACTCTGTCCTTGGAAGCACCTGCCAGAATTAAGAAC
3481 -----+-----+-----+-----+-----+-----+ 3540
CTCTCCTAGTTAGTGATGTAATTCTGAGACAGGAACCTTCGTGGACGGTCTTAATTCTTG

a E R I N H Y I K T L S L E A P A R I K N -

AAGGCTCCCTCCCCTGACTGGCCCCAGGAGGGAGAGGTGACCTTTGAGAACGCAGAGATG
3541 -----+-----+-----+-----+-----+-----+ 3600
TTCCGAGGGAGGGGACTGACCGGGGTCTCCCTCTCCACTGGAAACTCTTGCGTCTCTAC

a K A P S P D W P Q E G E V T F E N A E M -

AGGTACCGAGAAAACCTCCCTCTTGTCTAAAGAAAGTATCCTTCACGATCAAACCTAAA
3601 -----+-----+-----+-----+-----+-----+ 3660
TCCATGGCTCTTTTGGAGGGAGAACAGGATTTCTTTTCATAGGAAGTGCTAGTTTGGATTT

a R Y R E N L P L V L K K V S F T I K P K -

GAGAAGATTGGCATTGTGGGGCGGACAGGATCAGGGAAGTCCTCGCTGGGGATGGCCCTC
3661 -----+-----+-----+-----+-----+-----+ 3720
CTCTTCTAACCGTAACACCCCGCCTGTCTAGTCCCTTCAGGAGCGACCCCTACCGGGAG

a E K I G I V G R T G S G K S S L G M A L -

TTCCGTCTGGTGGAGTTATCTGGAGGCTGCATCAAGATTGATGGAGTGAGAATCAGTGAT
3721 -----+-----+-----+-----+-----+-----+ 3780
AAGGCAGACCACCTCAATAGACCTCCGACGTAGTTCTAACTACCTCACTCTTAGTCACTA

a F R L V E L S G G C I K I D G V R I S D -

ATTGGCCTTGCCGACCTCCGAAGCAAACCTCTCTATCATTCTCAAGAGCCGGTGCTGTTC
3781 -----+-----+-----+-----+-----+-----+ 3840
TAACCGGAACGGCTGGAGGCTTCGTTTGAGAGATAGTAAGGAGTTCTCGGCCACGACAAG

Figure 13I

33/56

a I G L A D L R S K L S I I P Q E P V L F -
AGTGGCACTGTCAGATCAAATTTGGACCCCTTCAACCAGTACACTGAAGACCAGATTTGG
3841 -----+-----+-----+-----+-----+-----+ 3900
TCACCGTGACAGTCTAGTTTAAACCTGGGGAAGTTGGTCATGTGACTTCTGGTCTAAACC

a S G T V R S N L D P F N O Y T E D O I W -
GATGCCCTGGAGAGGACACACATGAAAGAATGTATTGCTCAGCTACCTCTGAAACTTGAA
3901 -----+-----+-----+-----+-----+-----+ 3960
CTACGGGACCTCTCCTGTGTGTACTTTCTTACATAACGAGTCGATGGAGACTTTGAACTT

a D A L E R T H M K E C I A Q L P L K L E -
TCTGAAGTGATGGAGAATGGGGATAACTTCTCAGTGGGGGAACGGCAGCTCTTGTGCATA
3961 -----+-----+-----+-----+-----+-----+ 4020
AGACTTCACTACCTCTTACCCCTATTGAAGAGTCACCCCTTGCCGTCGAGAACACGTAT

a S E V M E N G D N F S V G E R Q L L C I -
GCTAGAGCCCTGCTCCGCACTGTAAGATTCTGATTTTAGATGAAGCCACAGCTGCCATG
4021 -----+-----+-----+-----+-----+-----+ 4080
CGATCTCGGGACGAGGCGGTGACATTCTAAGACTAAAATCTACTTCGGTGTGACGGTAC

a A R A L L R H C K I L I L D E A T A A M -
GACACAGAGACAGACTTATTGATTCAAGAGACCATCCGAGAAGCATTTGCAGACTGTACC
4081 -----+-----+-----+-----+-----+-----+ 4140
CTGTGTCTCTGTCTGAATAACTAAGTTCTCTGGTAGGCTCTTCGTAAACGTCTGACATGG

a D T E T D L L I Q E T I R E A F A D C T -
ATGCTGACCATTGCCCATCGCCTGCACACGGTTCTAGGCTCCGATAGGATTATGGTGCTG
4141 -----+-----+-----+-----+-----+-----+ 4200
TACGACTGGTAACGGGTAGCGGACGTGTGCCAAGATCCGAGGCTATCCTAATACCACGAC

a M L T I A H R L H T V L G S D R I M V L -
GCCAGGGACAGGTGGTGGAGTTTGACACCCCATCGGTCCTTCTGTCCAACGACAGTTCC
4201 -----+-----+-----+-----+-----+-----+ 4260
CGGGTCCCTGTCCACCACCTCAAACCTGTGGGGTAGCCAGGAAGACAGGTTGCTGTCAAGG

Figure 13J

34/56

a A Q G Q V V E F D T P S V L L S N D S S .

CGATTCTATGCCATGTTTGCTGCTGCAGAGAACAAGGTCGCTGTCAAGGGCTGA
4261 -----+-----+-----+-----+-----+----- 4314
GCTAAGATACGGTACAAACGACGACGTCTCTTGTTCAGCGACAGTTCCCGACT

a R F Y A M F A A A E N K V A V K G * .

Figure 13K

35/56

MOAT D cDNA AND AMINO ACID SEQUENCE ENCODED THEREBY

ATGGACGCCCTGTGCGGTTCCGGGGAGCTCGGCTCCAAGTTCTGGGACTCCAACCTGTCT
 1 -----+-----+-----+-----+-----+-----+ 60
 TACCTGCGGGACACGCCAAGGCCCTCGAGCCGAGGTTCAAGACCCTGAGGTTGGACAGA

a M D A L C G S G E L G S K F W D S N L S .

GTGCACACAGAAAACCCGGACCTCACTCCCTGCTTCCAGAACTCCCTGCTGGCCTGGGTG
 61 -----+-----+-----+-----+-----+-----+ 120
 CACGTGTGTCTTTTGGGCTGGAGTGAGGGACGAAGGTCTTGAGGGACGACCGGACCCAC

a V H T E N P D L T P C F Q N S L L A W V .

CCCTGCATCTACCTGTGGGTCGCCCTGCCCTGCTACTTGCTCTACCTGCGGCACCATTGT
 121 -----+-----+-----+-----+-----+-----+ 180
 GGGACGTAGATGGACACCCAGCGGGACGGGACGATGAACGAGATGGACGCCGTGGTAACA

a P C I Y L W V A L P C Y L L Y L R H H C .

CGTGGCTACATCATCCTCTCCACCTGTCCAAGCTCAAGATGGTCCTGGGTGTCCTGCTG
 181 -----+-----+-----+-----+-----+-----+ 240
 GCACCGATGTAGTAGGAGAGGGTGGACAGGTTGAGTTCTACCAGGACCCACAGGACGAC

a R G Y I I L S H L S K L K M V L G V L L .

TGGTGCGTCTCCTGGGCGGACCTTTTTTACTCCTTCCATGGCCTGGTCCATGGCCGGGCC
 241 -----+-----+-----+-----+-----+-----+ 300
 ACCACGCAGAGGACCCGCCTGGAAAAAATGAGGAAGGTACCGGACCAGGTACCGGCCCGG

a W C V S W A D L F Y S F H G L V H G R A .

CCTGCCCCTGTTTTCTTTGTCACCCCCTTGGTGGTGGGGGTACCATGCTGCTGGCCACC
 301 -----+-----+-----+-----+-----+-----+ 360
 GGACGGGGACAAAAGAAACAGTGGGGGAACCACCCCCAGTGGTACGACGACCGGTGG

a P A P V F F V T P L V V G V T M L L A T .

CTGCTGATACAGTATGAGCGGCTGCAGGGCGTACAGTCTTCGGGGGTCCTCATTATCTTC

Figure 14A

36/56

361 -----+-----+-----+-----+-----+-----+ 420
 GACGACTATGTCATACTCGCCGACGTCCCGCATGTCAGAAGCCCCCAGGAGTAATAGAAG

a L L I O Y E R L Q G V Q S S G V L I I F -

TGGTTCCTGTGTGTGGTCTGCGCCATCGTCCCATTCGCTCCAAGATCCTTTTAGCCAAG

421 -----+-----+-----+-----+-----+-----+ 480
 ACCAAGGACACACACCAGACGCGGTAGCAGGGTAAGGCGAGGTTCTAGGAAAATCGGTTC

a W F L C V V C A I V P F R S K I L L A K -

GCAGAGGGTGAGATCTCAGACCCCTTCGCTTCACCACCTTCTACATCCACTTTGCCCTG

481 -----+-----+-----+-----+-----+-----+ 540
 CGTCTCCCACTCTAGAGTCTGGGGAAGGCGAAGTGGTGGAAGATGTAGGTGAAACGGGAC

a A E G E I S D P F R F T T F Y I H F A L -

GTA CTCTCTGCCCTCATCTTGGCCTGCTTCAGGGAGAAACCTCCATTTTCTCCGCAAAG

541 -----+-----+-----+-----+-----+-----+ 600
 CATGAGAGACGGGAGTAGAACCGGACGAAGTCCCTCTTTGGAGGTAAAAAGAGGCGTTTC

a V L S A L I L A C F R E K P P F F S A K -

AATGTCGACCCTAACCCTACCTGAGACCAGCGCTGGCTTTCTCTCCCGCCTGTTTTTC

601 -----+-----+-----+-----+-----+-----+ 660
 TTACAGCTGGGATTGGGGATGGGACTCTGGTCGCGACCGAAAGAGAGGGCGGACAAAAAG

a N V D P N P Y P E T S A G F L S R L F F -

TGGTGGTTCACAAAGATGGCCATCTATGGCTACCGGCATCCCCTGGAGGAGAAGGACCTC

661 -----+-----+-----+-----+-----+-----+ 720
 ACCACCAAGTGTTTCTACCGGTAGATACCGATGGCCGTAGGGGACCTCCTCTTCTGGAG

a W W F T K M A I Y G Y R H P L E E K D L -

TGGTCCCTAAAGGAAGAGGACAGATCCCAGATGGTGGTGCAGCAGCTGCTGGAGGCATGG

721 -----+-----+-----+-----+-----+-----+ 780
 ACCAGGGATTTCCTTCTCCTGTCTAGGGTCTACCACCACGTCGTCGACGACCTCCGTACC

a W S L K E E D R S Q M V V Q Q L L E A W -

AGGAAGCAGGAAAAGCAGACGGCACGACACAAGGCTTCAGCAGCACCTGGGAAAAATGCC

781 -----+-----+-----+-----+-----+-----+ 840

Figure 14B

SUBSTITUTE SHEET (RULE 26)

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TCCTTCGTCCTTTTCGTCTGCCGTGCTGTGTTCCGAAGTCGTCGTGGACCCTTTTACGG

a R K Q E K Q T A R H K A S A A P G K N A -

TCCGGCGAGGACGAGGTGCTGCTGGGTGCCGGGCCAGGCCCGGAAGCCCTCCTTCCTG

841 ----- + ----- + ----- + ----- + ----- + ----- + 900

AGGCCGCTCCTGCTCCACGACGACCCACGGGCCGGTCCGGGGCCTTCGGGAGGAAGGAC

a S G E D E V L L G A R P R P R K P S F L -

AAGGCCCTGCTGGCCACCTTCGGCTCCAGCTTCCTCATCAGTGCCTGCTTCAAGCTTATC

901 ----- + ----- + ----- + ----- + ----- + ----- + 960

TTCCGGGACGACCGGTGGAAGCCGAGGTCAAGGAGTAGTCACGGACGAAGTTCGAATAG

a K A L L A T F G S S F L I S A C F K L I -

CAGGACCTGCTCTCCTTCATCAATCCACAGCTGCTCAGCATCCTGATCAGGTTTATCTCC

961 ----- + ----- + ----- + ----- + ----- + ----- + 1020

GTCCTGGACGAGAGGAAGTAGTTAGGTGTCGACGAGTCGTAGGACTAGTCCAAATAGAGG

a Q D L L S F I N P Q L L S I L I R F I S -

AACCCCATGGCCCCCTCCTGGTGGGGCTTCCTGGTGGCTGGGCTGATGTTCTGTGCTCC

1021 ----- + ----- + ----- + ----- + ----- + ----- + 1080

TTGGGGTACCGGGGAGGACCACCCCGAAGGACCACCGACCCGACTACAAGGACACGAGG

a N P M A P S W W G F L V A G L M F L C S -

ATGATGCAGTCGCTGATCTTACAACACTATTACCACTACATCTTTGTGACTGGGGTGAAG

1081 ----- + ----- + ----- + ----- + ----- + ----- + 1140

TACTACGTCAGCGACTAGAATGTTGTGATAATGGTGATGTAGAAACACTGACCCCACTTC

a M M Q S L I L Q H Y Y H Y I F V T G V K -

TTTCGTA CTGGGATCATGGGTGTCATCTACAGGAAGGCTCTGGTTATCACCAACTCAGTC

1141 ----- + ----- + ----- + ----- + ----- + ----- + 1200

AAAGCATGACCCTAGTACCCACAGTAGATGTCCTTCCGAGACCAATAGTGGTTGAGTCAG

a F R T G I M G V I Y R K A L V I T N S V -

AAACGTGCGTCCACTGTGGGGGAAATTGTCAACCTCATGTGAGTGGATGCCAGCGCTTC

1201 ----- + ----- + ----- + ----- + ----- + ----- + 1260

TTTGACGCAGGTGACACCCCTTTAACAGTTGGAGTACAGTCACCTACGGGTCGCGAAG

Figure 14C

38/56

a K R A S T V G E I V N L M S V D A Q R F .
ATGGACCTTGCCCCCTTCCTCAATCTGCTGTGGTCAGCACCCCTGCAGATCATCCTGGCG
1261 -----+-----+-----+-----+-----+-----+ 1320
TACCTGGAACGGGGGAAGGAGTTAGACGACACCAGTCGTGGGGACGTCTAGTAGGACCGC

a M D L A P F L N L L W S A P L Q I I L A .
ATCTACTTCCTCTGGCAGAACCTAGGTCCCTCTGTCTGGCTGGAGTCGCTTTTCATGGTC
1321 -----+-----+-----+-----+-----+-----+ 1380
TAGATGAAGGAGACCGTCTTGGATCCAGGGAGACAGGACCGACCTCAGCGAAAGTACCAG

a I Y F L W Q N L G P S V L A G V A F M V .
TTGCTGATTCCACTCAACGGAGCTGTGGCCGTGAAGATGCGCGCCTTCCAGGTAAAGCAA
1381 -----+-----+-----+-----+-----+-----+ 1440
AACGACTAAGGTGAGTTGCCTCGACACCGGCACTTCTACGCGCGGAAGGTCCATTTGTT

a L L I P L N G A V A V K M R A F Q V K Q .
ATGAAATTGAAGGACTCGCGCATCAAGCTGATGAGTGAGATCCTGAACGGCATCAAGGTG
1441 -----+-----+-----+-----+-----+-----+ 1500
TACTTTAACTTCCTGAGCGCGTAGTTGACTACTCACTCTAGGACTTGCCGTAGTTCCAC

a M K L K D S R I K L M S E I L N G I K V .
CTGAAGCTGTACGCCTGGGAGCCCAGCTTCCTGAAGCAGGTGGAGGGCATCCGGCAGGGT
1501 -----+-----+-----+-----+-----+-----+ 1560
GACTTCGACATGCGGACCCTCGGGTGAAGGACTTCGTCCACCTCCCGTAGGCCGTCCCA

a L K L Y A W E P S F L K Q V E G I R Q G .
GAGCTCCAGCTGCTGCGCACGGCGGCCTACCTCCACACCACAACCACCTTCACCTGGATG
1561 -----+-----+-----+-----+-----+-----+ 1620
CTCGAGGTGACGACGCGTGCCGCGGATGGAGGTGTGGTGTGGTGGGAAGTGGACCTAC

a E L Q L L R T A A Y L H T T T T F T W M .
TGCAGCCCCTTCCTGGTGACCCTGATCACCTCTGGGTGTACGTGTACGTGGACCCAAAC
1621 -----+-----+-----+-----+-----+-----+ 1680
ACGTCGGGGAAGGACCACTGGGACTAGTGGGAGACCCACATGCACATGCACCTGGGTTTG

Figure 14D

39/56

a C S P F L V T L I T L W V Y V Y V D P N -

AATGTGCTGGACGCCGAGAAGGCCTTTGTGTCTGTGTCCTTGTTTAATATCTTAAGACTT
 1681 -----+-----+-----+-----+-----+-----+ 1740
 TTACACGACCTGCGGCTCTTCCGGAAACACAGACACAGGAACAAATTATAGAATTCTGAA

a N V L D A E K A F V S V S L F N I L R L -

CCCCCAACATGCTGCCCCAGTTAATCAGCAACCTGACTCAGGCCAGTGTGTCTCTGAAA
 1741 -----+-----+-----+-----+-----+-----+ 1800
 GGGGAGTTGTACGACGGGGTCAATTAGTCGTTGGACTGAGTCCGGTCACACAGAGACTTT

a P L N M L P Q L I S N L T Q A S V S L K -

CGGATCCAGCAATTCCTGAGCCAAGAGGAACTTGACCCCCAGAGTGTGGAAAGAAAGACC
 1801 -----+-----+-----+-----+-----+-----+ 1860
 GCCTAGGTCGTTAAGGACTCGGTTCTCCTTGAAGTGGGGTCTCACACCTTTCTTTCTGG

a R I Q Q F L S Q E E L D P Q S V E R K T -

ATCTCCCCAGGCTATGCCATCACCATACACAGTGGCACCTTCACCTGGGCCCAGGACCTG
 1861 -----+-----+-----+-----+-----+-----+ 1920
 TAGAGGGGTCCGATACGGTAGTGGTATGTGTCACCGTGGAAGTGGACCCGGGTCCTGGAC

a I S P G Y A I T I H S G T F T W A Q D L -

CCCCCACTCTGCACAGCCTAGACATCCAGGTCCCGAAAGGGGCACTGGTGGCCGTGGTG
 1921 -----+-----+-----+-----+-----+-----+ 1980
 GGGGGGTGAGACGTGTGCGGATCTGTAGGTCCAGGGCTTTCCCCGTGACCACCGGCACCAC

a P P T L H S L D I Q V P K G A L V A V V -

GGGCCTGTGGGCTGTGGGAAGTCCTCCCTGGTGTCTGCCCTGCTGGGAGAGATGGAGAAG
 1981 -----+-----+-----+-----+-----+-----+ 2040
 CCCGGACACCCGACACCCTTCAGGAGGGACCACAGACGGGACGACCCTCTCTACCTCTTC

a G P V G C G K S S L V S A L L G E M E K -

CTAGAAGGCAAAGTGACATGAAGGCATGGATCCAGAACTGCACTCTTCAGGAAAACGTG
 2041 -----+-----+-----+-----+-----+-----+ 2100
 GATCTCCGTTTCACGTGTACTTCCGTACCTAGGTCTTGACGTGAGAAGTCCTTTTGCAC

a L E G K V H M K A W I Q N C T L Q E N V -

Figure 14E

SUBSTITUTE SHEET (RULE 26)

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CTTTTCGGCAAAGCCCTGAACCCCAAGCGCTACCAGCAGACTCTGGAGGCCTGTGCCTTG
2101 ----- + ----- + ----- + ----- + ----- + ----- + 2160
GAAAAGCCGTTTCGGGACTTGGGGTTCGCGATGGTCGTCTGAGACCTCCGGACACGGAAC

a L F G K A L N P K R Y Q Q T L E A C A L -

CTAGCTGACCTGGAGATGCTGCCTGGTGGGGATCAGACAGAGATTGGAGAGAAGGGCATT
2161 ----- + ----- + ----- + ----- + ----- + ----- + 2220
GATCGACTGGACCTCTACGACGGACCACCCCTAGTCTGTCTCTAACCTCTCTTCCCGTAA

a L A D L E M L P G G D Q T E I G E K G I -

AACCTGTCTGGGGGCCAGCGGCAGCGGGTCAGTCTGGCTCGAGCTGTTTACAGTGATGCC
2221 ----- + ----- + ----- + ----- + ----- + ----- + 2280
TTGGACAGACCCCCGGTCGCCGTCGCCAGTCAGACCGAGCTCGACAAATGTCACTACGG

a N L S G G Q R Q R V S L A R A V Y S D A -

GATATTTTCTTGCTGGATGACCCACTGTCCGCGGTGGACTCTCATGTGGCCAAGCACATC
2281 ----- + ----- + ----- + ----- + ----- + ----- + 2340
CTATAAAAGAACGACCTACTGGGTGACAGGCGCCACCTGAGAGTACACCGGTTCTGTAG

a D I F L L D D P L S A V D S H V A K H I -

TTTGACCACGTCATCGGGCCAGAAGGCGTGCTGGCAGGCAAGACGCGAGTGCTGGTGACG
2341 ----- + ----- + ----- + ----- + ----- + ----- + 2400
AAACTGGTGCACTAGCCCGGTCTTCCGCACGACCGTCCGTTCTGCGCTCACGACCACTGC

a F D H V I G P E G V L A G K T R V L V T -

CACGGCATTAGCTTCCTGCCCCAGACAGACTTCATCATTGTGCTAGCTGATGGACAGGTG
2401 ----- + ----- + ----- + ----- + ----- + ----- + 2460
GTGCCGTAATCGAAGGACGGGGTCTGTCTGAAGTAGTAACACGATCGACTACCTGTCCAC

a H G I S F L P Q T D F I I V L A D G Q V -

TCTGAGATGGGCCCCGTACCCAGCCCTGCTGCAGCGCAACGGCTCCTTTGCCAACTTTCTC
2461 ----- + ----- + ----- + ----- + ----- + ----- + 2520
AGACTCTACCCGGGCATGGGTCTGGGACGACGTCGCGTTGCCGAGGAAACGGTTGAAAGAG

a S E M G P Y P A L L Q R N G S F A N F L -

Figure 14F

SUBSTITUTE SHEET (RULE 26)

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TGCAACTATGCCCCGATGAGGACCAAGGGCACCTGGAGGACAGCTGGACCGCGTTGGAA
2521 -----+-----+-----+-----+-----+-----+ 2580
ACGTTGATACGGGGGCTACTCCTGGTTCCCGTGGACCTCCTGTGACCTGGCGCAACCTT

a C N Y A P D E D Q G H L E D S W T A L E -

GGTGCAGAGGATAAGGAGGCACTGCTGATTGAAGACACACTCAGCAACCACACGGATCTG
2581 -----+-----+-----+-----+-----+-----+ 2640
CCACGTCTCCTATTCTCCGTGACGACTAATTCTGTGTGAGTCGTTGGTGTGCCTAGAC

a G A E D K E A L L I E D T L S N H T D L -

ACAGACAATGATCCAGTCACCTATGTGGTCCAGAAGCAGTTTATGAGACAGCTGAGTGCC
2641 -----+-----+-----+-----+-----+-----+ 2700
TGTCTGTTACTAGGTCAGTGGATACACCAGGTCTTCGTCAAATACTCTGTGCACTCACGG

a T D N D P V T Y V V Q K Q F M R Q L S A -

CTGTCTCAGATGGGGAGGGACAGGGTCGGCCTGTACCCCGGAGGCACCTGGGTCCATCA
2701 -----+-----+-----+-----+-----+-----+ 2760
GACAGGAGTCTACCCCTCCCTGTCCCAGCCGGACATGGGGCCTCCGTGGACCCAGGTAGT

a L S S D G E G Q G R P V P R R H L G P S -

GAGAAGGTGCAGGTGACAGAGGCGAAGGCAGATGGGGCACTGACCCAGGAGGAGAAAGCA
2761 -----+-----+-----+-----+-----+-----+ 2820
CTCTTCACGTCCACTGTCTCCGCTTCGCTACCCCGTGACTGGGTCTCCTCTTTTCGT

a E K V Q V T E A K A D G A L T Q E E K A -

GCCATTGGCACTGTGGAGCTCAGTGTGTTCTGGGATTATGCCAAGGCCGTGGGGCTCTGT
2821 -----+-----+-----+-----+-----+-----+ 2880
CGGTAACCGTGACACCTCGAGTCACACAAGACCCTAATACGGTTCCGGCACCCCGAGACA

a A I G T V E L S V F W D Y A K A V G L C -

ACCACGCTGGCCATCTGTCTCCTGTATGTGGGTCAAAGTGCGGCTGCCATTGGAGCCAAT
2881 -----+-----+-----+-----+-----+-----+ 2940
TGGTGCGACCGGTAGACAGAGGACATACACCCAGTTTCACGCCGACGGTAACCTCGGTTA

a T T L A I C L L Y V G Q S A A A I G A N

GTGTGGCTCAGTGCCTGGACAAATGATGCCATGGCAGACAGTAGACAGAACAACACTTCC

Figure 14G

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2941 ----- + ----- + ----- + ----- + ----- + ----- + 3000
CACACCGAGTCACGGACCTGTTTACTACGGTACCGTCTGTCATCTGTCTTGTGGAAGG

a V W L S A W T N D A M A D S R Q N N T S -

CTGAGGCTGGGCGTCTATGCTGCTTTAGGAATTCTGCAAGGGTCTTGGTGATGCTGGCA

3001 ----- + ----- + ----- + ----- + ----- + ----- + 3060
GACTCCGACCCGAGATACGACGAAATCCTTAAGACGTTCCCAAGAACCACTACGACCGT

a L R L G V Y A A L G I L Q G F L V M L A -

GCCATGGCCATGGCAGCGGGTGGCATCCAGGCTGCCCGTGTGTTGCACCAGGCACTGCTG

3061 ----- + ----- + ----- + ----- + ----- + ----- + 3120
CGGTACCGGTACCGTCGCCCACCGTAGGTCCGACGGGCACACAACGTGGTCCGTGACGAC

a A M A M A A G G I Q A A R V L H Q A L L -

CACAACAAGATACGCTCGCCACAGTCCTTCTTTGACACCACACCATCAGGCCGCATCCTG

3121 ----- + ----- + ----- + ----- + ----- + ----- + 3180
GTGTTGTTCTATGCGAGCGGTGTCAGGAAGAACTGTGGTGTGGTAGTCCGGCGTAGGAC

a H N K I R S P Q S F F D T T P S G R I L -

AACTGCTTCTCCAAGGACATCTATGTCGTTGATGAGGTTCTGGCCCCTGTCATCCTCATG

3181 ----- + ----- + ----- + ----- + ----- + ----- + 3240
TTGACGAAGAGGTTCTGTAGATACAGCAACTACTCCAAGACCGGGGACAGTAGGAGTAC

a N C F S K D I Y V V D E V L A P V I L M -

CTGCTCAATTCCTTCTTCAACGCCATCTCCACTCTTGTGGTCATCATGGCCAGCACGCCG

3241 ----- + ----- + ----- + ----- + ----- + ----- + 3300
GACGAGTTAAGGAAGAAGTTGCGGTAGAGGTGAGAACACCAGTAGTACCGGTCGTGCGGC

a L L N S F F N A I S T L V V I M A S T P -

CTCTTCACTGTGGTCATCCTGCCCTGGCTGTGCTCTACACCTTAGTGACGCGCTTCTAT

3301 ----- + ----- + ----- + ----- + ----- + ----- + 3360
GAGAAGTGACACCAGTAGGACGGGGACCGACACGAGATGTGGAATCACGTCGCGAAGATA

a L F T V V I L P L A V L Y T L V Q R F Y -

GCAGCCACATCACGGCAACTGAAGCGGCTGGAATCAGTCAGCCGCTCACCTATCTACTCC

3361 ----- + ----- + ----- + ----- + ----- + ----- + 3420

Figure 14H

SUBSTITUTE SHEET (RULE 26)

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CGTCGGTGTAGTGCCGTTGACTTCGCCGACCTTAGTCAGTCGGCGAGTGGATAGATGAGG

a A A T S R O L K R L E S V S R S P I Y S -

CACTTTTCGGAGACAGTGACTGGTGCCAGTGTATCCGGGCCTACAACCGCAGCCGGGAT
3421 -----+-----+-----+-----+-----+ 3480
GTGAAAAGCCTCTGTCACTGACCACGGTCACAGTAGGCCCGGATGTTGGCGTCGGCCCTA

a H F S E T V T G A S V I R A Y N R S R D -

TTTGAGATCATCAGTGATACTAAGGTGGATGCCAACCAGAGAAGCTGCTACCCCTACATC
3481 -----+-----+-----+-----+-----+ 3540
AAACTCTAGTAGTCACTATGATTCCACCTACGGTTGGTCTCTTCGACGATGGGGATGTAG

a F E I I S D T K V D A N Q R S C Y P Y I -

ATCTCCAACCGGTGGCTGAGCATCGGAGTGGAGTTCGTGGGGAAGTGCCTGGTCTCTTT
3541 -----+-----+-----+-----+-----+ 3600
TAGAGGTTGGCCACCGACTCGTAGCCTCACCTCAAGCACCCCTTGACGCACCACGAGAAA

a I S N R W L S I G V E F V G N C V V L F -

GCTGCACTATTTGCCGTCATCGGGAGGAGCAGCCTGAACCCGGGGCTGGTGGGCCTTTCT
3601 -----+-----+-----+-----+-----+ 3660
CGACGTGATAAACGGCAGTAGCCCTCCTCGTCGGACTTGGGCCCCGACCACCCGAAAGA

a A A L F A V I G R S S L N P G L V G L S -

GTGTCCTACTCCTTGCAGGTGACATTTGCTCTGAAGTGGATGATACGAATGATGTCAGAT
3661 -----+-----+-----+-----+-----+ 3720
CACAGGATGAGGAACGTCCACTGTAAACGAGACTTGACCTACTATGCTTACTACAGTCTA

a V S Y S L Q V T F A L N W M I R M M S D -

TTGGAATCTAACATCGTGGCTGTGGAGAGGGTCAAGGAGTACTCCAAGACAGAGACAGAG
3721 -----+-----+-----+-----+-----+ 3780
AACCTTAGATTGTAGCACCGACACCTCTCCAGTTCCTCATGAGGTTCTGTCTCTGTCTC

a L E S N I V A V E R V K E Y S K T E T E -

GCGCCCTGGGTGGTGGGAAGGCAGCCGCCCTCCCGAAGGTTGGCCCCACGTGGGGAGGTG
3781 -----+-----+-----+-----+-----+ 3840
CGCGGGACCCACCACCTTCCGTGGCGGGAGGGCTTCCAACCGGGGGTGCACCCCTCCAC

Figure 14I

SUBSTITUTE SHEET (RULE 26)

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a A P W V V E G S R P P E G W P P R G E V -

GAGTTCCGGAATTATTCTGTGCGCTACCGGCCGGGCCTAGACCTGGTGCTGAGAGACCTG
 3841 -----+-----+-----+-----+-----+-----+ 3900
 CTCAAGGCCTTAATAAGACACGCGATGGCCGGCCCGGATCTGGACCACGACTCTCTGGAC

a E F R N Y S V R Y R P G L D L V L R D L -

AGTCTGCATGTGCACGGTGGCGAGAAGGTGGGGATCGTGGGCCGCACTGGGGCTGGCAAG
 3901 -----+-----+-----+-----+-----+-----+ 3960
 TCAGACGTACACGTGCCACCGCTCTTCCACCCCTAGCACCCGGCGTGACCCCGACCGTTC

a S L H V H G G E K V G I V G R T G A G K -

TCTTCCATGACCCTTTGCCTGTTCCGCATCCTGGAGGCGGCAAAGGGTGAAATCCGCATT
 3961 -----+-----+-----+-----+-----+-----+ 4020
 AGAAGGTACTGGGAAACGGACAAGGCGTAGGACCTCCGCCGTTTCCCACTTTAGGCGTAA

a S S M T L C L F R I L E A A K G E I R I -

GATGGCCTCAATGTGGCAGACATCGGCCTCCATGACCTGCGCTCTCAGCTGACCATCATC
 4021 -----+-----+-----+-----+-----+-----+ 4080
 CTACCGGAGTTACACCGTCTGTAGCCGGAGGTACTGGACGCGAGAGTCGACTGGTAGTAG

a D G L N V A D I G L H D L R S Q L T I I -

CCGCAGGACCCCATCCTGTTCTCGGGGACCCTGCGCATGAACCTGGACCCCTTCGGCAGC
 4081 -----+-----+-----+-----+-----+-----+ 4140
 GGCGTCCTGGGGTAGGACAAGAGCCCCTGGGACGCGTACTTGGACCTGGGGAAGCCGTCG

a P Q D P I L F S G T L R M N L D P F G S -

TACTCAGAGGAGGACATTTGGTGGGCTTTGGAGCTGTCCACCTGCACACGTTTGTGAGC
 4141 -----+-----+-----+-----+-----+-----+ 4200
 ATGAGTCTCCTCCTGTAAACCACCCGAAACCTCGACAGGGTGGACGTGTGCAAACACTCG

a Y S E E D I W W A L E L S H L H T F V S -

TCCCAGCCGGCAGGCCTGGACTTCCAGTGCTCAGAGGGCGGGGAGAATCTCAGCGTGGGC
 4201 -----+-----+-----+-----+-----+-----+ 4260
 AGGGTCGGCCGTCCGGACCTGAAGGTCACGAGTCTCCCGCCCCTCTTAGAGTCGCACCCG

Figure 14J

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a S Q P A G L D F Q C S E G G E N L S V G -

CAGAGGCAGCTCGTGTGCCTGGCCCGAGCCCTGCTCCGCAAGAGCCGCATCCTGGTTTTA
4261 -----+-----+-----+-----+-----+-----+ 4320
GTCTCCGTCGAGCACACGGACCGGGCTCGGGACGAGGCGTTCTCGGCGTAGGACCAAAAT

a Q R Q L V C L A R A L L R K S R I L V L -

GACGAGGCCACACCTGCCATCGACCTGGAGACTGACAACCTCATCCAGGCTACCATCCGC
4321 -----+-----+-----+-----+-----+-----+ 4380
CTGCTCCGGTGTGACGGTAGCTGGACCTCTGACTGTTGGAGTAGGTCCGATGGTAGGCG

a D E A T A A I D L E T D N L I Q A T I R -

ACCCAGTTTGATACCTGCACTGTCCTGACCATCGCACACCGGCTTAACACTATCATGGAC
4381 -----+-----+-----+-----+-----+-----+ 4440
TGGGTCAAACATGGACGTGACAGGACTGGTAGCGTGTGGCCGAATTGTGATAGTACCTG

a T Q F D T C T V L T I A H R L N T I M D -

TACACCAGGGTCCTGGTCCTGGACAAAGGAGTAGTAGCTGAATTTGATTCTCCAGCCAAC
4441 -----+-----+-----+-----+-----+-----+ 4500
ATGTGGTCCCAGGACCAGGACCTGTTTCCTCATCATCGACTTAAACTAAGAGGTGCGTTG

a Y T R V L V L D K G V V A E F D S P A N -

CTCATTGCAGCTAGAGGCATCTTCTACGGGATGGCCAGAGATGCTGGACTTGCCTAA
4501 -----+-----+-----+-----+-----+-----+ 4557
GAGTAACGTCGATCTCCGTAGAAGATGCCCTACCGGTCTCTACGACCTGAACGGATT

a L I A A R G I F Y G M A R D A G L A * -

Figure 14K

MOAT E cDNA AND AMINO ACID SEQUENCE ENCODED THEREBY

ATGCCGCGCCTGCTGAGCCCTGCGCGGGGAGGGGGTCTGGAACACAGACAGAGCCTGAA
1 ----- + ----- + ----- + ----- + ----- + ----- + 60
TACCGGCGCGGACGACTCGGGACGCGCCCCGTCCCCCAGACCTTGGTCTGTCTCGGACTT

a M A A P A E P C A G Q G V W N O T E P E .

CCTGCCGCCACCAGCCTGCTGAGCCTGTGCTTCCTGAGAACAGCAGGGGTCTGGGTACCC
61 ----- + ----- + ----- + ----- + ----- + ----- + 120
GGACGGCGGTGGTCTGGACGACTCGGACACGAAGGACTCTTGTCGTCCCCAGACCCATGGG

a P A A T S L L S L C F L R T A G V W V P .

CCCATGTACCTCTGGGTCTTGGTCCCCTACCTCCTCTTCATCCACCACCATGGCCGG
121 ----- + ----- + ----- + ----- + ----- + ----- + 180
GGGTACATGGAGACCCAGGAACCAGGGTAGATGGAGGAGAAGTAGGTGGTGGTACCGGCC

a P M Y L W V L G P I Y L L F I H H H G R .

GGCTACCTCCGGATGTCCCCACTCTTCAAAGCCAAGATGGTGCTTGGATTGCGCCTCATA
181 ----- + ----- + ----- + ----- + ----- + ----- + 240
CCGATGGAGGCCTACAGGGGTGAGAAGTTTCGGTTCTACCACGAACCTAAGCGGGAGTAT

a G Y L R M S P L F K A K M V L G F A L I .

GTCCTGTGTACCTCCAGCGTGGCTGTGCTCTTTGGAAAATCCAACAGGGAACGCCTGAG
241 ----- + ----- + ----- + ----- + ----- + ----- + 300
CAGGACACATGGAGGTCGCACCGACAGCGAGAAACCTTTTAGGTTGTCCCTTGCGGACTC

a V L C T S S V A V A L W K I Q Q G T P E .

GCCCCAGAATTCCTCATTCATCCTACTGTGTGGCTCACCACGATGAGCTTCGCAGTGTTT
301 ----- + ----- + ----- + ----- + ----- + ----- + 360
CGGGGTCTTAAGGAGTAAGTAGGATGACACACCGAGTGGTGCTACTCGAAGCGTCACAAG

a A P E F L I H P T V W L T T M S F A V F .

CTGATTCACACCGAGAGGAAAAAGGGAGTCCAGTCATCTGGAGTGCTGTTTGGTTACTGG
361 ----- + ----- + ----- + ----- + ----- + ----- + 420
GACTAAGTGTTGGCTCTCCTTTTCCCTCAGGTCAGTAGACCTCACGACAAACCAATGACC

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a L I H T E R K K G V Q S S G V L F G Y W -
CTTCTCTGCTTTGTCTTGCCAGCTACCAACGCTGCCAGCAGGCCTCCGGAGCGGGCTTC
421 -----+-----+-----+-----+-----+-----+ 480
GAAGAGACGAAACAGAACGGTCGATGGTTGCGACGGGTCGTCCGGAGGCCTCGCCCGAAG

a L L C F V L P A T N A A Q Q A S G A G F -
CAGAGCGACCCTGTCCGCCACCTGTCCACCTACCTATGCCTGTCTCTGGTGGTGGCACAG
481 -----+-----+-----+-----+-----+-----+ 540
GTCTCGCTGGGACAGGCGGTGGACAGGTGGATGGATACGGACAGAGACCACCACCGTGTC

a Q S D P V R H L S T Y L C L S L V V A Q -
TTTGTGCTGTCCTGCCTGGCGGATCAACCCCCCTTCTTCCCTGAAGACCCCCAGCAGTCT
541 -----+-----+-----+-----+-----+-----+ 600
AAACACGACAGGACGGACCGCCTAGTTGGGGGGAAGAAGGGACTTCTGGGGGTCGTCAGA

a F V L S C L A D Q P P F F P E D P Q Q S -
AACCCCTGTCCAGAGACTGGGGCAGCCTTCCCCTCCAAAGCCACGTTCTGGTGGGTTTCT
601 -----+-----+-----+-----+-----+-----+ 660
TTGGGGACAGGTCTCTGACCCCGTCGGAAGGGGAGGTTTCGGTGCAAGACCACCCAAAGA

a N P C P E T G A A F P S K A T F W W V S -
GGCCTGGTCTGGAGGGGATACAGGAGGCCACTGAGACCAAAGACCTCTGGTCGCTTGGG
661 -----+-----+-----+-----+-----+-----+ 720
CCGGACCAGACCTCCCCTATGTCCTCCGGTGACTCTGGTTTTCTGGAGACCAGCGAACCC

a G L V W R G Y R R P L R P K D L W S L G -
AGAGAAAACCTCCTCAGAAGAACTTGTTTCCCGGCTTGAAAAGGAGTGATGAGGAACCGC
721 -----+-----+-----+-----+-----+-----+ 780
TCTCTTTTGAGGAGTCTTCTGAACAAAGGGCCGAACCTTTTCTCACCTACTCCTTGGCG

a R E N S S E E L V S R L E K E W M R N R -
AGTGCAGCCCGGAGGCACAACAAGGCAATAGCATTTAAAGGAAAGGCGGCAGTGGCATG
781 -----+-----+-----+-----+-----+-----+ 840
TCACGTCGGGCCTCCGTGTTGTTCCGTTATCGTAAATTTTCTTTCCGCCGTCACCGTAC

Figure 15B

SUBSTITUTE SHEET (RULE 26)

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a S A A R R H N K A I A F K R K G G S G M .
AAGGCTCCAGAGACCGAGCCCTTCCTACGGCAAGAAGGGAGCCAGTGGCGCCCACTGCTG
841 -----+-----+-----+-----+-----+-----+ 900
TTCCGAGGTCTCTGGCTCGGGAAGGATGCCGTTCTTCCCTCGGTCACCGCGGGTGACGAC

a K A P E T E P F L R Q E G S Q W R P L L .
AAGGCCATCTGGCAGGTGTTCCATTCTACCTTCCTCCTGGGGACCCCTCAGCCTCATCATC
901 -----+-----+-----+-----+-----+-----+ 960
TTCCGGTAGACCGTCCACAAGGTAAGATGGAAGGAGGACCCCTGGGAGTCGGAGTAGTAG

a K A I W Q V F H S T F L L G T L S L I I .
AGTGATGTCTTCAGGTTCACTGTCCCCAAGCTGCTCAGCCTTTTCCTGGAGTTTATTGGT
961 -----+-----+-----+-----+-----+-----+ 1020
TCACTACAGAAGTCCAAGTGACAGGGGTTGACGAGTCGGAAAAGGACCTCAAATAACCA

a S D V F R F T V P K L L S L F L E F I G .
GATCCCAAGCCTCCAGCCTGGAAGGGCTACCTCCTCGCCGTGCTGATGTTCTCTCAGCC
1021 -----+-----+-----+-----+-----+-----+ 1080
CTAGGGTTTCGGAGGTCGGACCTTCCCGATGGAGGAGCGGCACGACTACAAGGAGAGTCGG

a D P K P P A W K G Y L L A V L M F L S A .
TGCCTGCAAACGCTGTTTGAGCAGCAGAACATGTACAGGCTCAAGGTGCCGCAGATGAGG
1081 -----+-----+-----+-----+-----+-----+ 1140
ACGGACGTTTTCGACAAACTCGTCGTCTTGACATGTCCGAGTTCCACGGCGTCTACTCC

a C L Q T L F E Q Q N M Y R L K V P Q M R .
TTGCGGTTCGGCCATCACTGGCCTGGTGTACAGAAAGGTCCTGGCTCTGTCCAGCGGCTCC
1141 -----+-----+-----+-----+-----+-----+ 1200
AACGCCAGCCGGTAGTGACCGGACCACATGTCTTCCAGGACCGAGACAGGTCGCCGAGG

a L R S A I T G L V Y R K V L A L S S G S .
AGAAAGGCCAGTGCGGTGGGTGATGTGGTCAATCTGGTGTCCGTGGACGTGCAGCGGCTG
1201 -----+-----+-----+-----+-----+-----+ 1260
TCTTTCGGTCACGCCACCCACTACACCAGTTAGACCACAGGCACCTGCACGTGCGCCGAC

a R K A S A V G D V V N L V S V D V Q R L .

Figure 15C

SUBSTITUTE SHEET (RULE 26)

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ACCGAGAGCGTCTCTACCTCAACGGGCTGTGGCTGCCTCTCGTCTGGATCGTGGTCTGC
1261 -----+-----+-----+-----+-----+-----+ 1320
TGGCTCTCGCAGGAGATGGAGTTGCCCGACACCGACGGAGAGCAGACCTAGCACCCAGACG

a T E S V L Y L N G L W L P L V W I V V C -

TTCTGTCTATCTCTGGCAGCTCCTGGGGCCCTCCGCCCTCACTGCCATCGCTGTCTTCCTG
1321 -----+-----+-----+-----+-----+-----+ 1380
AAGCAGATAGAGACCGTCGAGGACCCGGGAGGCGGGAGTGACGGTAGCGACAGAAGGAC

a F V Y L W Q L L G P S A L T A I A V F L -

AGCCTCCTCCCTCTGAATTTCTTCATCTCCAAGAAAAGGAACCACCATCAGGAGGAGCAA
1381 -----+-----+-----+-----+-----+-----+ 1440
TCGGAGGAGGGAGACTTAAAGAAGTAGAGGTTCTTTTCCTTGGTGGTAGTCCTCCTCGTT

a S L L P L N F F I S K K R N H H Q E E Q -

ATGAGGCAGAAGGACTCACGGGCACGGCTCACCAGCTCTATCCTCAGGAACCTCGAAGACC
1441 -----+-----+-----+-----+-----+-----+ 1500
TACTCCGTCTTCCTGAGTGCCCGTGCCGAGTGGTCGAGATAGGAGTCCTTGAGCTTCTGG

a M R Q K D S R A R L T S S I L R N S K T -

ATCAAGTTCCATGGCTGGGAGGGAGCCTTTCTGGACAGAGTCCTGGGCATCCGAGGCCAG
1501 -----+-----+-----+-----+-----+-----+ 1560
TAGTTCAAGGTACCGACCCTCCCTCGGAAAGACCTGTCTCAGGACCCGTAGGCTCCGGTC

a I K F H G W E G A F L D R V L G I R G Q -

GAGCTGGGCGCCTTGCGGACCTCCGGCCTCCTCTCTGTGTGCTGGTGTCTTCCAA
1561 -----+-----+-----+-----+-----+-----+ 1620
CTCGACCCGCGGAACGCCTGGAGGCCGGAGGAGAAGAGACACAGCGACCACAGGAAGGTT

a E L G A L R T S G L L F S V S L V S F Q -

GTGTCTACATTTCTGGTTCGCACTGGTGGTGGTTTGCTGTCCACACTCTGGTGGCCGAGAAT
1621 -----+-----+-----+-----+-----+-----+ 1680
CACAGATGTAAAGACCAGCGTGACCACCACAAACGACAGGTGTGAGACCACCGGCTCTTA

a V S T F L V A L V V F A V H T L V A E N -

Figure 15D

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GCTATGAATGCAGAGAAAGCCTTTGTGACTCTCACAGTTCTCAACATCCTCAACAAGGCC
1681 -----+-----+-----+-----+-----+-----+ 1740
CGATACTTACGTCTCTTTTCGGAAACACTGAGAGTGTCAAGAGTTGTAGGAGTTGTTCCGG

a A M N A E K A F V T L T V L N I L N K A -

CAGGCTTTCTGCCCCTTCTCCATCCACTCCCTCGTCCAGGCCCGGGTGTCTTTGACCGT
1741 -----+-----+-----+-----+-----+-----+ 1800
GTCCGAAAGGACGGGAAGAGGTAGGTGAGGGAGCAGGTCGCGGCCACAGGAAACTGGCA

a Q A F L P F S I H S L V Q A R V S F D R -

CTGGTCACCTTCCTCTGCCTGGAAGAAGTTGACCCTGGTGTCTGACTCAAGTTCCTCT
1801 -----+-----+-----+-----+-----+-----+ 1860
GACCAGTGGAAGGAGACGGACCTTCTTCAACTGGGACCACAGCATCTGAGTTCAAGGAGA

a L V T F L C L E E V D P G V V D S S S -

GGAAGCGCTGCCGGAAGGATTGCATCACCATACACAGTGCCACCTTCGCCTGGTCCCAG
1861 -----+-----+-----+-----+-----+-----+ 1920
CCTTCGCGACGGCCCTTCTAACGTAGTGGTATGTGTACGGTGGAAGCGGACCAGGGTC

a G S A A G K D C I T I H S A T F A W S Q -

GAAAGCCCTCCCTGCCTCCACAGAATAAACCTCACGGTGCCCCAGGGCTGTCTGCTGGCT
1921 -----+-----+-----+-----+-----+-----+ 1980
CTTTCGGGAGGGACGGAGGTGTCTTATTTGGAGTGCCACGGGGTCCCGACAGACGACCGA

a E S P P C L H R I N L T V P Q G C L L A -

GTTGTGGTCCAGTGGGGGCAGGGAAGTCTCCCTGCTGTCCGCCCTCCTTGGGGAGCTG
1981 -----+-----+-----+-----+-----+-----+ 2040
CAACAGCCAGGTCACCCCCGTCCCTTCAGGAGGGACGACAGGCGGGAGGAACCCCTCGAC

a V V G P V G A G K S S L L S A L L G E L -

TCAAAGGTGGAGGGGTTCTGTGAGCATCGAGGGTGTGTGGCCTACGTGCCCCAGGAGGCC
2041 -----+-----+-----+-----+-----+-----+ 2100
AGTTTCCACCTCCCCAAGCACTCGTAGCTCCACGACACCGGATGCACGGGGTCTCCGG

a S K V E G F V S I E G A V A Y V P Q E A -

TGGGTGCAGAACACCTCTGTGGTAGAGAATGTGTGCTTCGGGCAGGAGCTGGACCCACCC

Figure 15E

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2101 -----+-----+-----+-----+-----+-----+ 2160
ACCCACGTCTTGTGGAGACACCATCTCTTACACACGAAGCCCGTCCTCGACCTGGGTGGG
a W V Q N T S V V E N V C F G Q E L D P P .

TGGCTGGAGAGAGTACTAGAAGCCTGTGCCCTGCAGCCAGATGTGGACAGCTTCCCTGAG
2161 -----+-----+-----+-----+-----+-----+ 2220
ACCGACCTCTCTCATGATCTTCGGACACGGGACGTCGGTCTACACCTGTCTGAAGGGACTC
a W L E R V L E A C A L Q P D V D S F P E .

GGAATCCACACTTCAATTGGGGAGCAGGGCATGAATCTCTCCGGAGGCCAGAAGCAGCGG
2221 -----+-----+-----+-----+-----+-----+ 2280
CCTTAGGTGTGAAGTTAACCCTCGTCCCGTACTTAGAGAGGCCTCCGGTCTTCGTCGCC
a G I H T S I G E Q G M N L S G G Q K Q R .

CTGAGCCTGGCCCCGGGCTGTATACAGAAAGGCAGCTGTGTACCTGCTGGATGACCCCTG
2281 -----+-----+-----+-----+-----+-----+ 2340
GACTCGGACCGGGCCCCGACATATGTCTTCCGTCGACACATGGACGACCTACTGGGGGAC
a L S L A R A V Y R K A A V Y L L D D P L .

GCGGCCCTGGATGCCACGTTGGCCAGCATGTCTTCAACCAGGTCATTGGGCCTGGTGGG
2341 -----+-----+-----+-----+-----+-----+ 2400
CGCCGGGACCTACGGGTGCAACCGGTCGTACAGAAGTTGGTCCAGTAACCCGGACCACCC
a A A L D A H V G Q H V F N Q V I G P G G .

CTACTCCAGGGAACAACACGGATTCTCGTGACGCACGCACTCCACATCCTGCCCCAGGCT
2401 -----+-----+-----+-----+-----+-----+ 2460
GATGAGGTCCCTTGTGTGCCTAAGAGCACTGCGTGCGTGAGGTGTAGGACGGGGTCCGA
a L L Q G T T R I L V T H A L H I L P Q A .

GATTGGATCATAGTGCTGGCAAATGGGGCCATCGCAGAGATGGGTTCTACCAGGAGCTT
2461 -----+-----+-----+-----+-----+-----+ 2520
CTAACCTAGTATCACGACCGTTTACCCCGGTAGCGTCTCTACCCAAGGATGGTCTCGAA
a D W I I V L A N G A I A E M G S Y Q E L .

CTGCAGAGGAAGGGGGCCCTCGTGTGTCTTCTGGATCAAGCCAGACAGCCAGGAGATAGA
2521 -----+-----+-----+-----+-----+-----+ 2580

Figure 15F

SUBSTITUTE SHEET (RULE 26)

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GACGTCTCCTTCCCCGGGAGCACACAGAAGACETAGTTCGGTCTGTCGGTCCTCTATCT

a L Q R K G A L V C L L D Q A R Q P G D R -

GGAGAAGGAGAAACAGAACCTGGGACCAGCACCAAGGACCCAGAGGCACCTCTGCAGGC
 2581 -----+-----+-----+-----+-----+-----+ 2640
 CCTCTTCTCTTTGTCTTGACCTGGTCGTGGTTCCTGGGGTCTCCGTGGAGACGTCCG

a G E G E T E P G T S T K D P R G T S A G -

AGGAGGCCCGAGCTTAGACGCGAGAGGTCCATCAAGTCAGTCCCTGAGAAGGACCGTACC
 2641 -----+-----+-----+-----+-----+-----+ 2700
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a R R P E L R R E R S I K S V P E K D R T -

ACTTCAGAAGCCCAGACAGAGGTTCTCTGGATGACCCTGACAGGGCAGGATGGCCAGCA
 2701 -----+-----+-----+-----+-----+-----+ 2760
 TGAAGTCTTCGGGTCTGTCTCCAAGGAGACCTACTGGGACTGTCCCGTCTACCGGTCTGT

a T S E A Q T E V P L D D P D R A G W P A -

GGAAAGGACAGCATCCAATACGGCAGGGTGAAGGCCACAGTGCACCTGGCCTACCTGCGT
 2761 -----+-----+-----+-----+-----+-----+ 2820
 CCTTCTCTGTCTAGGTTATGCCGTCCCACTTCCGGTGTACGTGGACCGGATGGACGCA

a G K D S I Q Y G R V K A T V H L A Y L R -

GCCGTGGGCACCCCCCTCTGCCTCTACGCACTCTTCTCTTCTCTGCCAGCAAGTGGCC
 2821 -----+-----+-----+-----+-----+-----+ 2880
 CGGCACCCGTGGGGGGAGACGGAGATGCGTGAGAAGGAGAAGGAGACGGTCGTTACCGG

a A V G T P L C L Y A L F L F L C Q Q V A -

TCCTTCTGCCGGGGCTACTGGCTGAGCCTGTGGGCGGACGACCCTGCAGTAGGTGGGCAG
 2881 -----+-----+-----+-----+-----+-----+ 2940
 AGGAAGACGGCCCCGATGACCGACTCGGACACCCGCTGCTGGGACGTCATCCACCCGTC

a S F C R G Y W L S L W A D D P A V G G Q -

CAGACGCAGGCAGCCCTGCGTGGCGGGATCTTCGGGCTCCTCGGCTGTCTCCAAGCCATT
 2941 -----+-----+-----+-----+-----+-----+ 3000
 GTCTGCGTCCGTCCGGACGCACCGCCCTAGAAGCCCCGAGGAGCCGACAGAGGTTCCGGTAA

Figure 15G

SUBSTITUTE SHEET (RULE 26)

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a Q T Q A A L R G G I F G L L G C L O A I .
GGGCTGTTTGCCTCCATGGCTGCGGTGCTCCTAGGTGGGGCCCGGGCATCCAGGTTGCTC
3001 -----+-----+-----+-----+-----+-----+ 3060
CCCGACAAACGGAGGTACCGACGCCACGAGGATCCACCCCGGGCCCGTAGGTCCAACGAG

a G L F A S M A A V L L G G A R A S R L L .
TTCCAGAGGCTCCTGTGGGATGTGGTGCGATCTCCCATCAGCTTCTTTGAGCGGACACCC
3061 -----+-----+-----+-----+-----+-----+ 3120
AAGGTCTCCGAGGACACCCTACACCACGCTAGAGGGTAGTCGAAGAACTCGCCTGTGGG

a F Q R L L W D V V R S P I S F F E R T P .
ATTGGTCACCTGCTAAACCGCTTCTCCAAGGAGACAGACACGGTTGACGTGGACATTCCA
3121 -----+-----+-----+-----+-----+-----+ 3180
TAACCA GTGGACGATTTGGCGAAGAGGTTCTCTGTCTGTGCCAACTGCACCTGTAAGGT

a I G H L L N R F S K E T D T V D V D I P .
GACAAACTCCGGTCCCTGCTGATGTACGCCTTTGGACTCCTGGAGGTCAGCCTGGTGGTG
3181 -----+-----+-----+-----+-----+-----+ 3240
CTGTTTGAGGCCAGGGACGACTACATGCGGAAACCTGAGGACCTCCAGTCGGACCAACAC

a D K L R S L L M Y A F G L L E V S L V V .
GCAGTGGCTACCCCACTGGCCACTGTGGCCATCCTGCCACTGTTTCTCCTCTACGCTGGG
3241 -----+-----+-----+-----+-----+-----+ 3300
CGTCACCGATGGGGTGACCGGTGACACCGGTAGGACGGTGACAAAGAGGAGATGCGACCC

a A V A T P L A T V A I L P L F L L Y A G .
TTTCAGAGCCTGTATGTGGTTAGCTCATGCCAGCTGAGACGCTTGGAGTCAGCCAGCTAC
3301 -----+-----+-----+-----+-----+-----+ 3360
AAAGTCTCGGACATACACCAATCGAGTACGGTCGACTCTGCGAACCTCAGTCGGTCGATG

a F Q S L Y V V S S C Q L R R L E S A S Y .
TCGTCTGTCTGCTCCCACATGGCTGAGACGTTCCAGGGCAGCACAGTGGTCCGGGCATTC
3361 -----+-----+-----+-----+-----+-----+ 3420
AGCAGACAGACGAGGGTGTACCGACTCTGCAAGGTCCCGTCGTGTCAACAGGCCCGTAAG

Figure 15H

54/56

a S S V C S H M A E T F Q G S T V V R A F -

CGAACCCAGGCCCTCTTGTGGCTCAGAACAAATGCTCGCGTAGATGAAAGCCAGAGGATC
 3421 -----+-----+-----+-----+-----+-----+ 3480
 GCTTGGGTCCGGGGAGAACACCGAGTCTTGTTACGAGCGCATCTACTTTCGGTCTCCTAG

a R T Q A P L V A Q N N A R V D E S Q R I -

AGTTTCCCGCGACTGGTGGCTGACAGGTGGCTTGCGGCCAATGTGGAGCTCCTGGGGAAT
 3481 -----+-----+-----+-----+-----+-----+ 3540
 TCAAAGGGCGCTGACCACCGACTGTCCACCGAACGCCGTTACACCTCGAGGACCCCTTA

a S F P R L V A D R W L A A N V E L L G N -

GGCCTGGTGTTCGAGCTGCCACGTGTGCTGTGCTGAGCAAAGCCACCTCAGTGCTGGC
 3541 -----+-----+-----+-----+-----+-----+ 3600
 CCGGACCACAAACGTCGACGGTGCACACGACACGACTCGTTTCGGGTGGAGTCACGACCG

a G L V F A A A T C A V L S K A H L S A G -

CTCGTGGGCTTCTCTGTCTCTGCTGCCCTCCAGGTGACCCAGGCACTGCAGTGGGTTGTT
 3601 -----+-----+-----+-----+-----+-----+ 3660
 GAGCACCCGAAGAGACAGAGACGACGGGAGGTCCACTGGGTCCGTGACGTCACCCAACAA

a L V G F S V S A A L Q V T Q A L Q W V V -

CGCAACTGGACAGACCTAGAGAACAGCATCGTGTGAGTGGAGCGGATGCAGGACTATGCC
 3661 -----+-----+-----+-----+-----+-----+ 3720
 GCGTTGACCTGTCTGGATCTCTTGTCGTAGCACAGTCACCTCGCCTACGTCCTGATACGG

a R N W T D L E N S I V S V E R M Q D Y A -

TGGACGCCCAAGGAGGCTCCCTGGAGGCTGCCACATGTGCAGCTCAGCCCCCTGGCCT
 3721 -----+-----+-----+-----+-----+-----+ 3780
 ACCTGCGGGTTCCTCCGAGGGACCTCCGACGGGTGTACACGTCGAGTCGGGGGGACCGGA

a W T P K E A P W R L P T C A A Q P P W P -

CAGGGCGGGCAGATCGAGTTCGGGACTTTGGGCTAAGATACCGACCTGAGCTCCCGCTG
 3781 -----+-----+-----+-----+-----+-----+ 3840
 GTCCCGCCCGTCTAGCTCAAGGCCCTGAAACCCGATTCTATGGCTGGACTCGAGGGCGAC

a Q G G Q I E F R D F G L R Y R P E L P L -

Figure 15I

SUBSTITUTE SHEET (RULE 26)

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GCTGTGCAGGGCGTGTCCCTCAAGATCCACGCAGGAGAGAAGGTGGGCATCGTTGGCAGG
3841 -----+-----+-----+-----+-----+ 3900
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a A V Q G V S L K I H A G E K V G I V G R -

ACCGGGGCAGGGAAGTCCTCCCTGGCCAGTGGGCTGCTGCGGCTCCAGGAGGCAGCTGAG
3901 -----+-----+-----+-----+-----+ 3960
TGGCCCCGTCCCTTCAGGAGGGACCGGTCACCCGACGACGCCGAGGTCTCCGTCGACTC

a T G A G K S S L A S G L L R L Q E A A E -

GGTGGGATCTGGATCGACGGGGTCCCCATTGCCACGTGGGGCTGCACACACTGCGCTCC
3961 -----+-----+-----+-----+-----+ 4020
CCACCCTAGACCTAGCTGCCCCAGGGGTAACGGGTGCACCCCGACGTGTGTGACGCGAGG

a G G I W I D G V P I A H V G L H T L R S -

AGGATCAGCATCATCCCCAGGACCCCATCCTGTTCCCTGGCTCTCTGCGGATGAACCTC
4021 -----+-----+-----+-----+-----+ 4080
TCCTAGTCGTAGTAGGGGGTCTGGGGTAGGACAAGGGACCGAGAGACGCCTACTTGGAG

a R I S I I P Q D P I L F P G S L R M N L -

GACCTGCTGCAGGAGCACTCGGACGAGGCTATCTGGGCAGCCCTGGAGACGGTGCAGCTC
4081 -----+-----+-----+-----+-----+ 4140
CTGGACGACGTCCTCGTGAGCCTGCTCCGATAGACCCGTCGGGACCTCTGCCACGTCGAG

a D L L Q E H S D E A I W A A L E T V Q L -

AAAGCCTTGGTGGCCAGCCTGCCCGGCCAGCTGCAGTACAAGTGTGCTGACCGAGGCGAG
4141 -----+-----+-----+-----+-----+ 4200
TTTCGGAACCACCGGTCGGACGGGCGGTCGACGTCATGTTACACGACTGGCTCCGCTC

a K A L V A S L P G Q L Q Y K C A D R G E -

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4201 -----+-----+-----+-----+-----+ 4260
CTGGACTCGCACCCGGTCTTTGTGAGGACACAGACCGTGCACGGGAAGAGGCCTTCTGG

a D L S V G Q K Q L L C I A R A L L R K T -

Figure 15J

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CAGATCCTCATCCTGGACGAGGCTACTGCTGCCGTGGACCCTGGCACGGAGCTGCAGATG
4261 -----+-----+-----+-----+-----+-----+ 4320
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a Q I L I L D E A T A A V D P G T E L O M -

CAGGCCATGCTCGGGAGCTGGTTTGCACAGTGCACCTGTGCTGCTCATTGCCCACCGCCTG
4321 -----+-----+-----+-----+-----+-----+ 4380
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a Q A M L G S W F A Q C T V L L I A H R L -

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4381 -----+-----+-----+-----+-----+-----+ 4440
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a R S V M D C A R V L V M D K G Q V A E S -

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4441 -----+-----+-----+-----+-----+-----+ 4500
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a G S P A Q L L A Q K G L F Y R L A Q E S -

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a G L V * -

Figure 15K

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SEQUENCE LISTING

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Kruh, Gary D.
Lee, Kun
Belinsky, Martin G.
Bain, Lisa J.

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Nucleic Acids and Methods of Use Thereof

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Ser Gly Leu Thr Val Ala Thr Val Leu Phe Gly Ile Ala Arg Ser Leu						
	770			775		780
Leu Val Phe Tyr Val Leu Val Asn Ser Ser Gln Thr Leu His Asn Lys						
	785			790		800
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Pro Ile Gly Arg Ile Leu Asn Arg Phe Ser Lys Asp Ile Gly His Leu						
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Ser Arg Trp Phe Ala Val Arg Leu Asp Ala Ile Cys Ala Met Phe Val						
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 Asn Phe Ser Val Gly Glu Arg Gln Leu Leu Cys Ile Ala Arg Ala Leu
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5079

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 Leu Pro Cys Tyr Leu Leu Tyr Leu Arg His His Cys Arg Gly Tyr Ile
 50 55 60
 Ile Leu Ser His Leu Ser Lys Leu Lys Met Val Leu Gly Val Leu Leu
 65 70 75 80
 Trp Cys Val Ser Trp Ala Asp Leu Phe Tyr Ser Phe His Gly Leu Val
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 His Gly Arg Ala Pro Ala Pro Val Phe Phe Val Thr Pro Leu Val Val
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 Gly Val Thr Met Leu Leu Ala Thr Leu Leu Ile Gln Tyr Glu Arg Leu
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 Gln Gly Val Gln Ser Ser Gly Val Leu Ile Ile Phe Trp Phe Leu Cys
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 Val Val Cys Ala Ile Val Pro Phe Arg Ser Lys Ile Leu Leu Ala Lys
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 His Phe Ala Leu Val Leu Ser Ala Leu Ile Leu Ala Cys Phe Arg Glu
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 Lys Pro Pro Phe Phe Ser Ala Lys Asn Val Asp Pro Asn Pro Tyr Pro
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 Glu Thr Ser Val Gly Phe Leu Ser Arg Leu Phe Phe Trp Trp Phe Thr
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 Lys Met Ala Ile Tyr Gly Tyr Arg His Pro Leu Glu Glu Lys Asp Leu
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 Trp Ser Leu Lys Glu Asp Arg Ser Gln Met Val Val Gln Gln Leu
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 Leu Glu Ala Trp Arg Lys Gln Glu Lys Gln Thr Ala Arg His Lys Ala
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 Gly Ala Arg Pro Arg Pro Arg Lys Pro Ser Phe Leu Lys Ala Leu Leu
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 Gln Asp Leu Leu Ser Phe Ile Asn Pro Gln Leu Leu Ser Ile Leu Ile
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 His Tyr Tyr His Tyr Ile Phe Val Thr Gly Val Lys Phe Arg Thr Gly
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 Lys Arg Ala Ser Thr Val Gly Glu Ile Val Asn Leu Met Ser Val Asp
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 Ala Gln Arg Phe Met Asp Leu Ala Pro Phe Leu Asn Leu Leu Trp Ser
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 Ala Pro Leu Gln Ile Ile Leu Ala Ile Tyr Phe Leu Trp Gln Asn Leu
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Met	Lys	Leu	Lys	Asp	Ser	Arg	Ile	Lys	Leu	Met	Ser	Glu	Ile	Leu	Asn
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Gly	Ile	Lys	Val	Leu	Lys	Leu	Tyr	Ala	Trp	Glu	Pro	Ser	Phe	Leu	Lys
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Gln	Val	Glu	Gly	Ile	Arg	Gln	Gly	Glu	Leu	Gln	Leu	Leu	Arg	Thr	Ala
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Ala	Tyr	Leu	His	Thr	Thr	Thr	Thr	Phe	Thr	Trp	Met	Cys	Ser	Pro	Phe
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Leu	Val	Thr	Leu	Ile	Thr	Leu	Trp	Val	Tyr	Val	Tyr	Val	Asp	Pro	Asn
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Asn	Val	Leu	Asp	Ala	Glu	Lys	Ala	Phe	Val	Ser	Val	Ser	Leu	Phe	Asn
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Ile	Leu	Arg	Leu	Pro	Leu	Asn	Met	Leu	Pro	Gln	Leu	Ile	Ser	Asn	Leu
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Thr	Gln	Ala	Ser	Val	Ser	Leu	Lys	Arg	Ile	Gln	Gln	Phe	Leu	Ser	Gln
	595						600					605			
Glu	Glu	Leu	Asp	Pro	Gln	Ser	Val	Glu	Arg	Lys	Thr	Ile	Ser	Pro	Gly
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Pro	Pro	Thr	Leu	His	Ser	Leu	Asp	Ile	Gln	Val	Pro	Lys	Gly	Ala	Leu
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Ala	Leu	Leu	Gly	Glu	Met	Glu	Lys	Leu	Glu	Gly	Lys	Val	His	Met	Lys
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Gln	Gln	Thr	Leu	Glu	Ala	Cys	Ala	Leu	Leu	Ala	Asp	Leu	Glu	Met	Leu
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Pro	Gly	Gly	Asp	Gln	Thr	Glu	Ile	Gly	Glu	Lys	Gly	Ile	Asn	Leu	Ser
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Gly	Gly	Gln	Arg	Gln	Arg	Val	Ser	Leu	Ala	Arg	Ala	Val	Tyr	Ser	Asp
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Ala	Asp	Ile	Phe	Leu	Leu	Asp	Pro	Leu	Ser	Ala	Val	Asp	Ser	His	
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Val	Ala	Lys	His	Ile	Phe	Asp	His	Val	Ile	Gly	Pro	Glu	Gly	Val	Leu
785				790					795						800
Ala	Gly	Lys	Thr	Arg	Val	Leu	Val	Thr	His	Gly	Ile	Ser	Phe	Leu	Pro
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Gln	Thr	Asp	Phe	Ile	Ile	Val	Leu	Ala	Asp	Gly	Gln	Val	Ser	Glu	Met
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Gly	Pro	Tyr	Pro	Ala	Leu	Leu	Gln	Arg	Asn	Gly	Ser	Phe	Ala	Asn	Phe
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Leu	Cys	Asn	Tyr	Ala	Pro	Asp	Glu	Asp	Gln	Gly	His	Leu	Glu	Asp	Ser
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Asp	Thr	Leu	Ser	Asn	His	Thr	Asp	Leu	Thr	Asp	Asn	Asp	Pro	Val	Thr
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Asp	Tyr	Ala	Lys	Ala	Val	Gly	Leu	Cys	Thr	Thr	Leu	Ala	Ile	Cys	Leu
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 995 1000 1005
 Ser Leu Arg Leu Gly Val Tyr Ala Ala Leu Gly Ile Leu Gln Gly Phe
 1010 1015 1020
 Leu Val Met Leu Ala Ala Met Ala Met Ala Ala Gly Gly Ile Gln Ala
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 Ala Arg Val Leu His Gln Ala Leu Leu His Asn Lys Ile Arg Ser Pro
 1045 1050 1055
 Gln Ser Phe Phe Asp Thr Thr Pro Ser Gly Arg Ile Leu Asn Cys Phe
 1060 1065 1070
 Ser Lys Asp Ile Tyr Val Val Asp Glu Val Leu Ala Pro Val Ile Leu
 1075 1080 1085
 Met Leu Leu Asn Ser Phe Phe Asn Ala Ile Ser Thr Leu Val Val Ile
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 Lys Arg Leu Glu Ser Val Ser Arg Ser Pro Ile Tyr Ser His Phe Ser
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 Glu Thr Val Thr Gly Ala Ser Val Ile Arg Ala Tyr Asn Arg Ser Arg
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 1475 1480 1485
 Asp Tyr Thr Arg Val Leu Val Leu Asp Lys Gly Val Val Ala Glu Phe
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INTERNATIONAL SEARCH REPORT

International application No.
PCT/US99/06644

A. CLASSIFICATION OF SUBJECT MATTER

IPC(6) : A01N 63/00, A61K 39/395, C12N 15/00, A01N 61/00, C07H 21/02
US CL : Please See Extra Sheet.

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 424/93.1, 93.2, 130.1; 435/320.1, 325; 514/1; 536/23.1; 800/13, 18

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

APS, STN, MEDLINE, BIOSIS, CAPLUS, SCISEARCH

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	Database GENBANK, Accession No. U66687, ALLIKMETS et al. Characterization of the human ABC superfamily: isolation and mapping of 21 new genes using the expressed sequence tags databases. Hum. Mol. Genet. 5(10), pp. 1649-1655, 26 March 1997.	21
X	Database GENBANK, Accession No. D77412, NISHIGUCHI. S. et al., A catalogue of genes in mouse embryonal carcinoma F9 cells identified with expressed sequence tags. J. Biochem. 119 (4), pp. 749-767, 04 October 1996.	22

☒ Further documents are listed in the continuation of Box C. ☐ See patent family annex.

A	document defining the general state of the art which is not considered to be of particular relevance	*T*	later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
E	earlier document published on or after the international filing date	*X*	document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
L	document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)	*Y*	document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
O	document referring to an oral disclosure, use, exhibition or other means	*Z*	document member of the same patent family
P	document published prior to the international filing date but later than the priority date claimed		

Date of the actual completion of the international search

20 MAY 1999

Date of mailing of the international search report

01 JUL 1999

Name and mailing address of the ISA/US
Commissioner of Patents and Trademarks
Box PCT
Washington, D.C. 20231

Facsimile No. (703) 305-3230

Authorized officer

SHIN-LIN CHEN

Telephone No. (703) 308-0196

INTERNATIONAL SEARCH REPORT

International application No.
PCT/US99/06644

C (Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	Database GENBANK, Accession No. U66674, ALLIKMETS, R. et al., Characterization of the human ABC superfamily: isolation and mapping of 21 new genes using the expressed sequence tags database. Hum. Mol. Genet. 16 March 1997, 5 (10), pp. 1649-1655.	33
X	Database GENBANK, Accession No. R97754, HILLIER, L. et al., The WashU-Merk EST project. 11 September 1995.	44
Y	KOIKE et al. A Canalicular Multispecific Organic Anion Transporter (cMOAT) Antisense cDNA Enhances Drug Sensitivity in Human Hepatic Cancer Cells. Cancer Research. 15 December 1997, Vol. 57, No. 24, pages 5475-5479, see entire document.	55-57
A,P	LEE et al. Isolation of MOAT-B, a Widely Expressed Multidrug Resistance-associated Proteins Canalicular Multispecific Organic Anion Transporter-related Transporter. Cancer Research. 01 July 1998, Vol 58, No. 13, pages 2741-2747, see entire document.	1-58
A,P	BELINSKY et al. Characterization of MOAT-C and MOAT-D, New Members of the MRP/cMOAT Subfamily of Transporter Proteins. Natl. Cancer Inst. 18 November 1998, Vol 90, No. 22, pages 1735-1741.	1-58
A	SUZUKI et al. Excretion of GSSG and Glutathione Conjugates Mediated by MRP1 and cMOAT/MRPS. Seminars in Liver Disease. 1998, Vol 18, No. 4, pages 359-376.	1-58

INTERNATIONAL SEARCH REPORT

International application No.
PCT/US99/06644

A. CLASSIFICATION OF SUBJECT MATTER:

US CL :

424/93.1, 93.2, 130.1; 435/320.1, 325; 514/1; 536/23.1; 800/13, 18